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Setching solution for etching porous silicon, etching method using the etching solution and method of preparing semiconductor member using the etching solution.

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67 A method for preparing a semiconductor member comprises: forming a substrate having a non-porous silicon monocrystalline layer and a porous silicon layer; bonding another substrate having a surface made of an insulating material to the surface of the monocrystalline layer; and exching to remove the porous silicon layer by immersing in an tohing solution.

## BACKGROUND OF THE INVENTION

## Fild of th Inv ntion

This inv ntion relates to an etching solution for porous silicon, an tiching method using the etching is lution and a method of producing a semiconductor memb r using the toling solution. Particularly, this invention relates to a method of producing a semiconductor member which is suitable for separation of dielectric materials or electronic devices, integrated circuits prepared on a monocrystalline semiconductor layer on an insulating material.

## Related Background Art

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Formation of a monocrystalline SI semiconductor layer on an insulating material has been widely known as the silicon on insulator (SOI) technology, and since a large number of advantages which cannot be reached by bulk SI substrates for preparation of conventional SI integrated circuits are possessed by the device utilizing the SOI structure, so many researches have been done. More specifically, by utilizing the SOI structure, the following advantages can be obtained:

- 1. Dielectric isolation can be easily done to enable high degree of integration;
- Radiation hardness is excellent;
  - 3. Stray capacity is reduced to attain high speed;
  - 4. Well formation step can be omitted;
  - 5. Latch-up can be prevented;
  - 5. Later-up can be prevented,
  - 6. Fully depleted field effect transistor can be made by thin film formation.

In order to realize the many advantages in device characteristics as mentioned above, studies have been made about the method for forming the SOI structure for these some 10 years. The contents are summarized in, for example, the literature as mentioned below:

Special Issue: "Single-rystal silicon on non-single-crystal insulators"; edited by G. W. Cullen, Journal of Crystal Growth, Volume 63, No. 3, pp. 429 - 590 (1963).

Also, it has been known for a long time to form the SOS (silicon on sapphire) structure by heteroepitaxy of sion a monocrystalline sapphire substrate by CVD (chemical vapor deposition) method. This was successful to some extent as the most mature SOI technique, but for such reasons as a large amount of crystal defects because of lattice mismatching at the Interface between the Si layer and the sapphire substrate, Introduction of aluminium from the sapphire substrate into the Si layer, and above all the high cost of the substrate and delay in enlargement of the substrate wafer size, it is obstructed from being widely applied. Relatively in recent years, attempts to realize the SOI structure without use of a sapphire substrate have been done. Such attempts may be broadly classified into the two shown below:

- (1) After surface exidation of an Si monocrystalline substrate, a window is formed to have the Si substrate partially exposed, and epitaxial growth is proceeded in the lateral direction with that exposed portion as the seed to form an Si monocrystalline layer on SiO<sub>2</sub>. (In this case, deposition of Si layer on SiO<sub>2</sub> is accompanied).
- (2) By use of an Si monocrystalline substrate itself as an active layer, SiO<sub>2</sub> is formed therebeneath. (This method is accompanied wit no deposition of SI layer).

As the means for realizing the above (1), there have been known the method in which a monocrystalline SI isyer is formed directly to lateral epitaxial growth by CVD, the method in which amorphous SI is deposited and subjected to solid phase lateral epitaxial growth by heat treatment, the method in which an amorphous or polycrystalline Isi Isi grown on SIO, by melting and recrystallization, and the method in which a melting region is scanned in a zone fashion by a rod-shaped heater (Zone meiting recrystallization). These methods have both advantages and disadvantages, they still have many problems with respect to controllability, productivity, uniformity and quality, and none of them have been industrially applied yet up to date. For example, the CVD method requires sacrifice-oxidation in flat thin film formation, while the crystallinity is poor in the solid phase growth method. On the other hand, in the been annealing method, problems are involved in controllability such as treatment time by converged beam scanning, the manner of overlapping of beams, focus adjustment, etc. Among these, the Zone Melting Recrystallization method is the most mature, and a relatively larger scale integrated circuit has been trially made, but still a large number of crystal defects such as point defects, line defects, plane defects (sub-boundary), etc. remain, and no device driven by minority carriers has been prepared.

Concerning the m thod using no Si substrat as th seed for epitaxial growth which is th ab v method

(2), for example, the following method may be included.

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Or

- 1. An oxid film is formed on an SI monocrystallin substrate with V-grooves as anis tropically tched on th surface, a polycrystallin SI lay risd posited on the oxid film thick to th extent as th SI substrate, and th reafter by polishing from the back surface of the Si substrate, Si monocrystalline regions di lectrically separated by surrounding with th V-grooves on th thick polycrystalline SI lay r are formed. In this method, although crystallinity is good, there are problems with respect to controllability and productivity in the step of depositing the polycrystalline SI thick as some hundred microns and the step in which the monothestep of depositing the polycrystalline SI thick as some hundred microns and the step in which the monothestep of the polycrystalline SI thick as some hundred microns and the step in which the monothestep of the polycrystalline SI thick as some hundred microns and the step in which the monothestep of the polycrystalline SI thick as some hundred microns and the step in which the monothestep of the polycrystalline SI thick as some hundred microns and the step in which the monothestep of the polycrystalline SI thick as some hundred microns and the step in which the monothestep of the polycrystalline SI thick as some hundred microns and the step in which the monothestep of the polycrystalline SI thick as some hundred microns and the step in which the monothestep of the polycrystalline SI thick as some hundred microns and the step in the polycrystalline SI thick as some hundred microns and the step in the step crystalline SI substrate is polished from the back surface to leave only the SI active layer as separated.
- 2. This is the method called SIMOX (Separation by ion-implanted oxygen) in which an SiO<sub>2</sub> layer is formed by ion implantation of oxygen into an SI monocrystalline substrate, which is one of the most mature methods because of good matching with the SHC (Integrated Circuit) process. However, for formation of the SIO<sub>2</sub> layer, 1018 ions/cm² or more of oxygen ions are required to be implanted, and the implantation time is very long to be not high in productivity, and also the wafer cost is high. Further, many crystal defects remain, and from an industrial point of view, no sufficient level of quality capable of preparing a device driven by
- 3. This is the method to form an SOI structure by dielectric isolation according to oxidation of porous SI. This is a method in which an N-type SI layer is formed on the surface of a P-type SI monocrystalline substrate in shape of islands by way of proton ion implantation (Imal et al., J. Crystal Growth, Vol. 63, 547 (1983)), or by epitaxial growth and patterning; only the P-type Si substrate is made porous by anodization in HF solution so as to surround the Si islands from the surface; and then the N-type Si islands are dielectrically Isolated by accelerated oxidation. In this method, the separated SI region is determined before the device steps, whereby there is the problem that the degree of freedom in drive and circuit design may be

A light-transmissive substrate is important for forming a contact sensor serving as a light-receiving device and a projection-type liquid crystal image display. A high-quality driving device is required for further increasing the density, resolution and definition of the pixels (picture element) of such a sensor or display. It is consequently necessary to produce a device to be provided on a light-transmissive substrate by using a monocrystalline layer

However, if an Si layer is deposited on a light-transmissive substrate such as glass substrate, etc., the Si having excellent crystallinity. layer is generally an amorphous layer or, at best, a polycrystalline layer because the Si layer reflects the disorder of the crystal structure of the substrate, and no high-quality device can thus be formed by using the Si layer. This is because the substrate has an amorphous crystal structure, and thus a monocrystalline layer of high quality cannot be easily obtained by simply depositing the Si layer, it is therefore difficult to produce a driving device having properties sufficient for the present demands or future demands because the crystal structure of an amorphous Si or polycrystalline Si has many defects.

Further, any one of the methods using an SI monocrystalline substrate is unsuitable for obtaining a good

monocrystalline film on a light-transmissive substrate. Takao Yonehara, one of the inventors, previously proposed a method of forming a semiconductor substrate

which is capable of solving the above problems in Japanese Patent Application No. 2-206548. The method of forming a semiconductor substrate disclosed in Patent Application No. 2-205548 comprises forming a substrate having a non-porous semiconductor monocrystalline layer and a porous semiconductor layer, bonding another substrate having an insulating material surface to the surface of the monocrystalline

layer, and removing the porous semiconductor layer by etching. This Invention has been achieved for improving the Invention disclosed in Patent Application No. 2-206548

The method of forming a semiconductor substrate disclosed in Patent Application No. 2-206548 comprises previously proposed. the step of removing porous SI by selective etching.

Porous SI was discovered in the course of research on electrolytic polishing of a semiconductor which was

conducted by Uhilir et al, in 1956 (A. Uhilir, Bell Syst. Tech. J., Vol. 35, pp 333 (1956)). Unagami et al. Investigated dissolving reaction of SI during anodization and reported that the anodic reaction of SI in a HF solution requires positive holes, and that the reaction is expressed as follows (T. Unagami,

tion of Si in a Hr solution required (1980)); J. Electrochem. Soc., Vol. 127, pp 476 (1980)); 
$$SI+2HF+(2-n)e^+ \rightarrow SIF_2+2H^+ + ne^-$$
 (1)  $SIF_2+2HF \rightarrow SIF_4+H_2$  (2)  $SIF_4+2HF \rightarrow H_2SIF_6$  (3)

$$SI + 4HF + (4-\lambda)^+ \rightarrow SiF_4 + 4H^+ + \lambda e^-$$
 (4)

$$SIF_4 + 2HF \rightarrow H_2SiF_6$$
 (5)

wherein <sup>+</sup> and <sup>-</sup> respectiv ly denote a positive hole and an lectron, and n and λ each d notes th number of positive hol a required for dissolving one allicon atom. Porous SI can be formed when the condition, n > 2 or λ > 4, is satisfied.

It is therefore found that positive holes are required for forming porous Si, and that P-type Si can be more easily made porous than N-type SI. However, it is also known that N-type SI can be made porous if holes are implanted thereto (R. P. Holmstrom and J. Y. Chi, Appl. Phys. Lett., Vol. 42, 386 (1983)).

The density of the porous SI layer can be changed to the range of 1.1 to 0.8 g/cm<sup>2</sup> by changing the concentration of the HF solution from S0 to 20%, as compared with the density of 2.33 g/cm<sup>2</sup> of monocrystalline SI. The porous SI layer has pores having an average size of about 600 Å which was measured by observation by a transmission electron microscope. Although the porous SI layer has a density which is half or less than that of monocrystalline SI, monocrystallinity is maintained, and a monocrystalline SI layer can be formed on the porous Is aver by ealthast growth.

Although the volume of an SI monocrystal is generally increased by 2.2 times by oxidation, the increase is in volume can be suppressed by controlling the density of the porous SI so that the occurrence of curvature of a substrate or the occurrence of a crack in a monocrystalline layer remained on the surface can be avoided during the oxidation process. The volume ratio R of monocrystalline SI to porous SI after oxidation can be expressed as follows:

$$R = 2.2 \times (A/2.33)$$
 (6)

wherein A denotes the density of porous Si. If R = 1, i.e., there is no increase in volume after oxidation, A = 1.08 (g/cm²). Namely, if the density of the porous Si layer is 1.08, an increase in volume, which is caused by oxidation, can be suppressed.

It can be said that at present, porous SI is subjected as such directly to subsequent steps (epitaxial growth and oxidation) after producting it, and the porous Si itself is not processed. This is because the porous Si cannot be easily processed or removed with good controllability. Namely, it has been not reported yet that porous SI is eitched with good controllability.

In addition, P generally shown by the following equation is referred as porosity:

$$P = (2.33 - A)/2.33$$
 (7)

When the value of porosity is adjusted to 30 to 55% during anodization, the properties of oxidized porous Si can be equalized to those of a monocrystalline Si oxide film. The porosity is expressed as follows:

$$P = (m1 - m2)/(m1 - m3)$$
 (8)

or

$$P = (m1 - m2)/pAt$$
 (9)

wherein

35 m1: total weight before anodization

m2: total weight after anodization

m3: total weight after removal of porous SI

density of monocrystalline Si

A: area of porous region

t thickness of porous SI

ceeds as follows:

However, the area of the porous region cannot be accurately calculated in many cases. In this case, although the equation (8) is effective, the porous SI must be etched for measuring the value of m3.

In addition, during epitaxial growth on the porous Si, the porous Si is capable of relieving distortion produced during heteroepitaxial growth and suppressing the occurrence of defects. However, in this case, since it is clear that the prossity is a very important parameter, measurement of the porosity is necessary and Indispensable. Known methods of etching porous Si are the following methods (1) and (2):

- (1) The method of etching porous SI with an aqueous NaOH solution (G. Bonchil, R. Herino, K. Barla, and J. C. Pfister, J. Electrochem. Soc., Vol. 130, No. 7, 1611 (1983)).
- (2) The method of etching porous SI with an etching solution which is capable of etching non-porous SI. In the above method (2), a fluoronitric acid-type etching solution is generally used, and etching of SI pro-

$$SI + 2O \rightarrow SIO_2$$
 (10)  
 $SIO_2 + 4HF \rightarrow SiF_4 + H_2O$  (11)

As shown by the above reaction formulas, Si is oxidized to SiO<sub>2</sub>, and the SiO<sub>2</sub> produced is etched with hydrofluoric acid.

Examples of etching solutions for non-porous SI include the above fluoronitric acid-type etching solution as well as ethylenediamine-type, KOH-type and hydrazine-type etching solutions and the like.

In this invention, it is necessary in selective tching of porous Si to select an etching solution which is cap-

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able of etching porous SI, other than th above tching solutions for non-porous SI. The porous SI is generally selectively etch d by the above method (1) which us an aqueous NaOH solution as an tching solution.

As d scrib d abov, both porous and non-porous Si are etched with the fluoronitric acid-type etchings is

On the other hand, in the conventional mithod of selectivity itching porous Si with an aqueous NaOH obution, Na ions are inevitably adsorbed on the etched surface. Since the Na ions cause impurity contamination, are movable and have adverse effects such as the formation of a interfacial states, the ions must not be introduced into the semiconductor process.

## 10 SUMMARY OF THE INVENTION

An object of this invention is to provide an etching solution which efficiently, uniformly, selectively and chemically etches porous SI without affecting the semiconductor process and etching non-porous SI.

Another object of this invention is to provide a method of preparing a semiconductor member using an etching solution for uniformly and selectively etching porous Si.

## BRIEF DESCRIPTION OF THE DRAWINGS

Figs. 1A and 1B are schematic views explaining an etching step using an etching solution of this invention;

Figs. 2A and 2 Bare schematic views explaining an etching step using an etching solution of this invention; Figs. 2A and 2B are schematic views explaining an etching step using an etching solution of this invention; Figs. 3A to 3C are schematic views explaining an etching step using an etching solution of this invention; Figs. 5A to 5D are schematic views explaining an etching step using an etching solution of this invention; Figs. 5A to 5D are schematic views explaining an etching step using an etching solution of this invention;

Figs. 6A to 6H are graphs showing the etching properties of porous and non-porous Si when etching solutions of this invention are respectively used;

Figs. 7A to 7H are graphs showing the relations between the etched thickness (etching depth) of porous SI and etching time when etching solutions of this invention are respectively used; Figs. 8A to 8C are schematic views explaining a process for preparing a semiconductor member of this

invention;
Figs. 9A to 9D are schematic views explaining a process for preparing a semiconductor member of this

rigs. 94 to 90 are scrientatic views explaining a process for preparing a semiconductor member of this Figs. 10A to 10C are schematic views explaining a process for preparing a semiconductor member of this

invention;

Figs. 11A to 11D are schematic views explaining a process for preparing a semiconductor member of this invention:

Figs. 12A to 12C are schematic views explaining a process for preparing a semiconductor member of this invention:

Figs. 13A to 13C are schematic views explaining a process for preparing a semiconductor member of this invention; and

Figs. 14A to 14D are schematic views explaining a process for preparing a semiconductor member of this invention

# DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

According to one aspect of this invention, this invention provides a chemical etching solution for etching

A first embodiment of the chemical etching solution of this invention is hydrofluoric acid.

A second embodiment of the chemical etching solution is a mixture containing hydrofluonic acid and an alcohol

A third embodiment of the chemical etching solution is a mixture containing hydrofluoric acid and hydrogen peroxide.

A fourth embodiment of the chemical etching solution is a mixture containing hydrofluoric acid, an alcohol and hydrogen peroxide.

A fifth embodiment of the chemical etching solution is buffered hydrofluoric acid.

55 A sixth embodiment of the chemical etching solution is a mixture containing buffered hydrofluoric acid and an alcohol

A seventh embodiment of the chemical etching solution is a mixture containing buffered hydrofluoric acid and hydrogen peroxid.

A eighth imbodiment of thi chemical etching solution is a mixture of buffered hydrofluoric acid, an alcohol and hydrogen peroxide.

The tching method of this invention comprises selectively tching porous silicon using the tching solution of this Inv ntion.

According to anoth ir asplict of this invintion, this invention provides a mithod of preparing a semiconduc-

A first embodiment of the method of preparing a semiconductor member of this invention comprises forming a substrate having a non-porous monocrystalline silicon layer and a porous silicon layer, bonding another substrate having a surface made of insulating material to the surface of the monocrystalline layer, and etching the porous silicon layer by immersing it in hydrofluoric acid.

A second embodiment of the method of preparing a semiconductor member of this invention uses each of the second to eighth forms of the etching solution of this invention in place of hydrofluoric acid used as an etching solution in the first form of the method of preparing a semiconductor member of this invention.

A third embodiment of the method of:preparing a semiconductor member of this invention comprises the steps of making a silicon substrate porous, forming a non-porous monocrystalline silicon layer on the silicon substrate made porous, bonding a light-transmissive glass substrate to the surface of the non-porous monocrystalline silicon layer, and selectively etching porous silicon so as to remove porous silicon by chemical etching using an etching solution of this invention by immersing the silicon substrate made porous therein.

A fourth embodiment of the method of preparing a semiconductor member of this invention comprises the steps of making a silicon substrate porous, forming a non-porous monocrystalline silicon layer on the silicon substrate made porous, bonding another silicon substrate having an insulating layer on the surface thereof to the surface of the non-porous monocrystalline silicon layer, and selectively etching porous silicon so as to remove porous silicon by chemical etching using an etching solution of this invention by immersing the silicon substrate made porous therein.

A fifth embodiment of the method of preparing a semiconductor member of this invention comprises the steps of making a silicon substrate porous, forming a non-porous monocrystalline silicon layer on the silicon substrate made porous, forming an oxide layer on the surface of the non-porous monocrystalline silicon layer, bonding a light-transmissive substrate to the surface of the oxide layer and selectively etching the silicon substrate made porous to remove it by chemical etching using an etching solution of this invention by immersing 30 the silicon substrate made porous therein.

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A sixth embodiment of the method of preparing a semiconductor member of this invention comprises the steps of making a silicon substrate porous, forming a non-porous monocrystalline silicon layer on the silicon substrate made porous, forming an oxide layer on the surface of the non-porous monocrystalline silicon layer. bonding another silicon substrate having an insulating layer on the surface thereof to the oxide layer formed on the non-porous monocrystalline silicon layer, and selectively etching the silicon substrate made porous to remove it by chemical etching using an etching solution of this invention by immersing the silicon substrate made porous therein.

In each of the above embodiments according to the method of preparing a semiconductor member of the present invention, the etching step may be carried out with coating the surfaces other than the surface of the silicon layer made porous with a protecting material before etching.

The etching solution for porous SI of this Invention is capable of uniformly and efficiently etching porous Si without the danger of contaminating the semiconductor process.

The etching method of this invention can be applied to usual semiconductor processes and is capable of selectively etching, with high accuracy, the porous SI provided on the same substrate provided non-porous SI to remove the porous SI because a chemical etching solution which does not etch non-porous SI is used.

The method of preparing a semiconductor member of this invention is excellent in productivity, uniformity, controllability and economy for forming a crystalline SI layer having excellent crystallinity equal to that of a monocrystalline wafer on insulating substrates such as light-transmissive insulating substrates represented by a glass substrate.

The method of preparing a semiconductor member of this invention is capable of realizing the advantages of conventional SOI devices and can be applied to various fields.

The method of preparing a semiconductor member of this invention can also be used in place of the expensive SOS or SIMOX used for producing a large-scale integrated circuit having the SOI structure.

In addition, the method of preparing a semiconductor member of this invention comprises the steps of chemically removing the lower portion of a monocrystalline SI substrate of high quality used as a starting matenal, with leaving only a monocrystalline layer on the surface thereof, and bonding the substrate to an insulating layer, and thus enables many treatments to be performed for a short time and has excellent productivity and economy.

Further, the method of preparing a semiconductor member of this inventi in can use a chi mical interest that it is the control of the control solution which has a bad eff ct on th s miconductor process in tching f porou SI and xhibits an etching selection ratio of a fiv digits value or more of porous SI to non-porous SI and xcellent controllability and productivity.

I. A description will now be given of the etching solution in accordance with the present invention.

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A description will be made first as to the case where hydrofluoric acid is used as the electroless wet chemical etching solution for porous SI, with specific reference to Fig. 7A.

Fig. 7A shows the etching time dependency of etched thickness of porous SI when the latter is etched by being immersed in hydrofluoric acid. The porous SI was formed by anodizing a monocrystalline SI. The conditions of anodization are shown below. It is to be noted, however, that the starting material for producing porous Si by anodization is not limited to monocrystalline SI and SI of other crystalline structure may be used as the starting material.

2.6 (V) Voltage applied: 30 (mA·cm<sup>-2</sup>)

Current density: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1 Anodizing solution:

2.4 hours Time 300 (um)

Thickness of porous Si: 56 (%)

Porosity:

Test pieces of the porous Si thus prepared were immersed in 49 % hydrofluoric acid solution (white circles) and 20 % hydrofluoric acid solution (black circles) both at the room temperature, and the solutions were agitated. The reduction in the thickness of these test pieces of the porous SI were then measured. A large etching rate was observed; namely, 90 μm in 40 minutes in case of the 49 % hydro hydro acid solution and 72 μm in 40 minutes in case of the 20 % hydrofluoric acid solution. After elapse of 80 minutes, the porous SI test pieces were etched by 205 μm with a high degree of surface state in case of the 49 % hydrofluoric acid solution, whereas, in case of the 20 % hydrofluoric acid solution, the etching depth was 164 µm after elapse of 80 minutes.

The etching rate has dependencies on the concentration of the etching solution and the temperature. The etching solution concentration and temperature are suitably determined in practical ranges. Although solution concentration of 49 % and 20 % and the room temperature are mentioned above, these are only illustrative and are not intended for limiting purpose. Preferably, the concentration of the hydrofluoric acid ranges between 5 % and 95 % and the temperature of the solution is set to a level which is ordinarily adopted in this field of technology.

The etched test places of porous Si were then rinsed with water and the surfaces after the rinsing were examined by microanalysis using secondary ions but no impurity was detected.

A description will now be given with specific reference to Fig. 6A of the etching characteristics of porous Si and non-porous SI when hydrofluonic acid is used as the etching solution, followed by a description of an example of etching of a non-porous SI substrate whose one side surface is completely porous Si, taken in conjunction with Figs. 1A and 1B.

Fig. 6A is a graph showing the time dependency of etching depth of porous SI and monocrystalline SI as observed when they are immersed in hydrofluoric acid solution. Porous SI was formed by anodizing monocrystalline Si under the same conditions as those mentioned above. In this case also, the use of monocrystalline Si as the starting material for anodization is only illustrative and Si of other crystalline structures may be used as the starting material.

Test pieces of the porous Si thus prepared were immersed in 49 % hydrofluoric acid solution (white circles) at the room temperature, and the solutions were agitated. The reduction in the thickness of these test pieces of porous SI were then measured. A large etching rate was observed: namely, 90 µm in 40 minutes in case of the 49 % hydrofluoric acid solution and, after elapse of 80 minutes, the porous SI test pieces were etched by  $205\ \mu m$  with a high degree of surface state. The etching rate has dependencies on the concentration of the etching solution and the temperature. The etching solution concentration and temperature are suitably determined in practical ranges. Although solution density of 49 % and the room temperature are mentioned above, these are only liustrative and are not intended for limiting purpose. Preferably, the concentration of the hydroflu ric acid ranges between 5 % and 95 % and the temperature of the solution is set to a level which is ordnarily adopt d in this field of t chn logy.

A test piece of a non-porous SI of 500 µm thick was immersed in a 49 % solution of hydrofluoric acid (black

circles), followed by an agitation of the solution. The reduction in the thickn as was thin measured. In this case, the test pile of non-porous SI was etched only by 100 Angstrom in riess even after eleps of 120 minutes. The tohing rate showed almost noid pendency on a lution concentration and it imperature.

Both th porous and non-porous Si test pieces aft r th etching were rinsed with water and th surface states of thes test pi ces wer xamin d by microanalysis with s condary I ns but no impurity was detected.

As shown in Fig. 1A, a monocrystallin SI substrate 22 was anodized nly at its ne sid s as to have a porous Si structure only at its one side as denoted by 21. Then, the substrate having the porous Si monocrystalline SI structure was Immersed in a hydrofluoric acid. As a consequence, only the porous SI portion was removed by the etching while the monocrystalline SI substrate 22 alone remained unetched, it is thus possible to selectively etch porous SI by using monocrystalline SI as the etch stopper.

A description will now be given of a case where both porous SI portion and monocrystalline SI portion are provided on one side of the substrate.

As shown in Fig. 2A, a portion of one side of a monocrystalline SI substrate 32 was anodized to become porous SI structure 31. Since the current and voltage necessary for the anodization vary depending on the carrier concentration, it is possible to selectively form porous SI structure by locally varying the carrier concentration, it is possible to selectively form porous SI structure by locally varying the carrier concentration, it in possible to selectively form porous SI structure was then immersed in hydrofluoric acid. As a result, only the porous SI/monocrystalline SI structure was then immersed in hydrofluoric acid. As a result, only the porous SI portion was removed while the monocrystalline SI substrate 32 remained unetched. It is thus possible to selectively etch porous SI.

A description will be made as to the case where a porous SI structure and a polycrystalline structure are formed in layers on one side of the substrate.

As shown in Fig. 3A, a polycrystalline SI layer 41 was formed by deposition on a single-crystalline SI substrate 42, and the surface layer of this polycrystalline Si was changed into a porous SI layer 43 by anodization, as shown in Fig. 3B. The substrate having the porous SI/polycrystalline SI/monocrystalline SI structure was immersed in a solution of hydrofluoric acid, whereby the porous SI structure alone was removed by etching while the monocrystalline SI substrate 42 and the polycrystalline SI layer 41 remained unched. It was thus possible to selectively etch the porous SI by using polycrystalline SI as the etching stopper.

A description will now be given of a case where porous Si portion is formed in the surface of a polycrystalline Si layer which is formed on one side of the substrate.

As shown in Fig. 4A, a polycrystalline Si layer 51 was formed by deposition in a monocrystalline SI substrate 52 and a portion of this polycrystalline Si layer was changed into porous Si layer 53 by anodization. Then, the 52 by a portion of this polycrystalline Si monocrystalline Si structure was immersed in a southor hydrofluoric acid, so that the porous Si elone was removed while the monocrystalline Si substrate 52 and the polycrystalline Si layer 51 remained unetched. It was thus possible to selectively etch the porous Si

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A description will now be given of the case where a mixture of hydrofluoric acid and an alcohol is used as the electroless wet chemical etching solution for porous Si, with reference to Fig. 7B.

Fig. 7B shows the time dependency of etching thickness of porous SI as observed when the porous SI is Immersed in a mixture liquid of hydrofluoric acid and ethyl alcohol without agitation of the liquid. The porous SI was formed by anodizing monocrystalline SI under the conditions shown below. The use of the monocrystalline SI as the starting material for forming the porous SI structure through anodization is only illustrative and SI of other crystalline structures can be used as the starting material.

5 Voltage applied: 2.6 (V)
Current density: 30 (mA-cm<sup>-2</sup>)
Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>3</sub>OH = 1:1:1
Time: 2.4 (hours)

Time: 2.4 (nours)
Thickness of porous Si: 300 (μm)
Percetty: 56 (%)

Porosity: 55 (%)

Test places of the porous SI prepared as described above were immersed, without agitation, in a mixture solution of 49 % hydrofluoric acid and ethyl alcohol (10 : 1) (white circles) and in a mixture solution of 20 % hydrofluoric acid and ethyl alcohol (10 : 1) (black circles). The reductions in the thicknesses of the porous SI hydrofluoric acid and ethyl alcohol (10 : 1) (black circles). The reductions in the thicknesses of the porous SI were observed: namely, in the case test places were then measured. Large rates of etching of the porous SI were observed: namely, in the case side and etching of the porous SI was etched by SS pum

of the mixture solution of 49 % hydrofluoric acid and ethyl alcohol (10:1), the porous Si was etched by 85 µm and, in case of the mixture solution of 20 % hydrofluoric acid and ethyl alcohol (10:1), the porous Si was etched by 68 µm, in about 40 minutes. After elaps of 80 minutes, the prous Si was etch d by a thickness as large as 195 µm in the cas of the mixtur solution of 49 % hydroflu ric acid and ethyl alcohol I (10:1) and 158 µm

even in th case of th mixture solution of 20 % hydrofluoric acid and ethyl alcoh 1 (10:1), with high degrees of states of the tohed surfaces.

The tching rate has depend noise on the concentration of the hydrofluoric acid solution, as will as in the temp rature. This addition of alcoholis rives to remove bubbles of reaction product gases ginerated as a result of this etching without delay from this surface billing etched, without necessitating agitation, thus insuring a high efficiency and uniformity of this etching.

The solution concentration and the temperature are determined such that a practical etching speed is obtained in preparation process and the effect of addition of alcohol is appreciable. Although the imbuture solutions of 49 % hydrofluoric acid and ethyl alcohol (10:1), as well as the room temperature as the solution temperature, are mentioned, these solution densities and temperature are only illustrative and are not hieraded to restrict the scope of the invention. The HF concentration with respect to the etching solution preferably ranges between 1 and 95 %, more preferably between 5 and 80 %. The concentration of alcohol with respect to the etching solution is preferably 60 % or less and most preferably 40 % or less, and is determined so as to provide an appreciable effect of addition of the alcohol. The temperature is selected to range preferably to 100°C, more preferably 56 80°C and most preferably 56 80°C.

Although ethyl alcohol has been mentioned specifically, the invention does not exclude the use of other alcohol such as isopropyl alcohol which does not cause any inconvenience in the production process and which can provide an appreciable effect of addition of such alcohol.

The porous Si after the etching was rinsed with water and the rinsed surface was examined by microanalysis by using secondary ions but no impurity was detected.

A description will now be given of the etching characteristic of porous SI and non-porous SI when they are etched by a mixture solution of hydrofluoric acid and ethyl alcohol, with specific reference to Fig. 8B.

Fig. 6B shows time dependencies of etched thicknesses of porous SI and monocrystalline SI as observed when the porous SI and the monocrystalline SI are immersed in a mixture solution of hydrofluoric acid and ethyl alcohol without agitation. The porous SI was formed by anodization of monocrystalline SI conducted under the same conditions as those shown before. The use of monocrystalline SI as the starting material for forming porous SI through anodization is only illustrative and SI of other crystalline structures can be used as the starting

A test piece of porous SI prepared as described above was immersed, without agitation, in a mixture solution of 49 % hydrofluoric acid and ethyl alcohol(10 : 1) (while circles), and reduction in the thickness of the porous SI was measured. The porous SI was repidly etched: namely, by a thickness of 85 μm in 40 minutes and 195 μm in 80 minutes, with high degrees of surface quality and uniformity.

The etching rate depends on the concentration and the temperature of the hydrofluoric acid solution.

The addition of alcohol serves to remove bubbles of reaction product gases generated as a result of the etching without delay from the surface being etched, without necessitating agitation, thus ensuring a high efficiency and uniformity of the etching.

A test piece of a non-porous Si of 500 µm thick was immersed in a mixture solution of 49 % hydrofluoric acid and ethyl alcohol (10:1) (black circles), without agitation of the solution. The reduction in the thickness was then measured. In this case, the test piece of non-porous Si was etched only by 100 Angstrom or less even after elapse of 120 minutes. The etching rate showed almost no dependency on the solution concentration and temperature.

Both the porous and non-porous SI test pieces after the etching were rinsed with water and the surface states of these test pieces were examined by microanalysis with secondary lone but no impurity was detected.

Naturally, various etching methods explained in connection with I by making reference to Figs. 1A and 1B, Figs. 2A and 2B, Figs. 3A to 3C and Figs. 4A to 4C can be realized also in the case where the mixture solution of hydrofluoric acid and an alcohol is used as the etchant for porous SI.

## H(3)

A description will now be given of the case where a mixture of hydrofluoric acid and aqueous hydrogen peroxide (hereinafter also referred to as "hydrogen peroxide") is used as the electroless wet chemical etching solution for porous SI, with reference to Fig. 7C.

Fig. 7C shows the time dependency of etched thickness of porous Si as observed when the porous Si is immersed in a mixture liquid of hydrofluoric acid and hydrogen peroxide under aglitation of the liquid. The porous Si was formed by anodizing monocrystalline Si under the conditions shown below. The use of the monocrystalline Si as the starting material for forming the porous Si structure through anodization is nly illustrative and Si of other crystalline structures can be used as the starting material.

Voltage applied: 2.6 (V)

Current density: 30 (mA·cm<sup>-2</sup>)

Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

Time: 2.4 (hours)
Thickn ss of porous Si: 300 (µm)
Porceity: 56 (%)

20

25

50

Porosity: 56 (%)

Test pleces of the porous SI prepared as described above were immersed, without agitation, in a mixture solution of 49 % hydrofluoric acid and hydrogen peroxide (1:5) (white circles) and in a mixture solution of 49 % hydrofluoric acid and hydrogen peroxide (1:1) (black circles). The reductions in the thitchnesses of the porous SI test pleces were then measured. Large rates of etching of the porous SI were observed: namely, in the case of the 1:5 solution, the porous SI was etched by 115 µm and, in case of the 1:1 solution, the porous SI was etched by 135 µm, in about 40 minutes. After elapse of 60 minutas, the porous SI was etched by a thickness alarge as 25 µm in the case of the 1:5 solution with high degrees of states of the etched surfaces. The concentration and 307 µm in the case of the 1:1 solution, with high degrees of states of the etched surfaces. The concentration of hydrogen peroxide was 30 % in this case but the hydrogen peroxide concentration may be determined in a range which provides an appreciable effect of addition of hydrogen peroxide and which does not cause any practical problem in the production process.

The etching rate has dependencies on the density of the hydrofluoric acid solution, as well as on the temperature of the same. The addition of alcohol serves to accelerate oxidation of silicon, thus enhancing the reaction speed as compared to the case where hydrogen peroxide is not used. It is also possible to control the reaction speed by suitably selecting the content of hydrogen peroxide.

The solution concentration and the solution temperature are determined such that a practical etching speed is obtained in preparation process and the effect of hydrofluoric acid and hydrogen peroxide is appreciable. Although the mixture solutions of 49 % hydrofluoric acid and hydrogen proxide (1:5) and 49 % hydrofluoric acid and hydrogen peroxide (1:1), as well as the room temperature as the solution temperature, are mentioned, these solution concentrations and temperature are only illustrative and are not intended to restrict the south

The HF concentration with respect to the etching solution preferably ranges between 1 and 95 %, more preferably between 5 and 90 % and most preferably between 5 and 80 %. The concentration of H<sub>2</sub>D, with respect to the etching solution is preferably selected to range between 1 and 95 %, more preferably between 5 and 90% and most preferably between 10 and 80%, and is determined so as to provide an appreciable effect of addition of the hydrogen peroxide. The temperature is selected to range preferably 0 to 100°C, more preferably 5 to 80°C and most preferably 5 to 80°C.

The porous SI after the etching was rinsed with water and the rinsed surface was examined by microanalysis by using secondary ions but no impurity was detected.

A description will now be given of the etching characteristics of porous Si and non-porous Si as observed when they are etched by a mixture solution of hydrofluoric acid and aqueous hydrogen peroxide, with specific reference to Fig. 6C. Fig. 6C shows the time dependencies of etched depth of porous Si and monocrystalline Si as observed when the porous Si and the monocrystalline Si are immersed in a mixture solution of hydrofluoric acid and aqueous hydrogen peroxide. The porous Si was formed by anodization of monocrystalline Si conducated and aqueous hydrogen peroxide. The porous Si was formed by anodization of monocrystalline Si as the starting material for forming porous Si through anodization is only illustrative and Si of other crystalline structures can be used as the starting material.

A test piece of porous SI prepared as described above was immersed, followed by agitation, in a mixture solution of 49 % hydrofluoric acid and aqueous hydrogen peroxide (white circles), and reduction in the thickness of the porous SI was measured. The porous SI was rapidly etched: namely, by a thickness of 112 µm in 40 minutes and 256 µm in 80 minutes, with high degrees of surface quality and uniformity. Although the concentration of aqueous hydrogen peroxide was 30 % in this case, the concentration of hydrogen peroxide may be suitably determined within a range which does not cause any practical inconvenience in the production.

The etching rate depends on the concentration and the temperature of the hydrofluoric acid and aqueous hydrogen peroxide

The addition of hydrogen peroxide serves to accelerate oxidation of silicon, thus enhancing the reaction speed as compared to the case where hydrogen peroxide is not added. Furthermore, the reaction speed can be controlled by suitably selecting the content of the hydrogen peroxide.

A test piece of a non-porous SI of 500 µm thick was immersed in a mixture solution of 49 % hydrofluoric acid and aqueous hydrogen peroxide (1:5) (Black circles), followed by agitation of the solution. The reduction in th thickn ss of th porous SI was then measured. In thickn say the test piece of nonporous SI was etched only by 100 Angstrom or I se even after elaps of 120 minutes. The etching rate showed aimost no dep indency on the solution concentration and temperature.

Both the porous and non-porous SI test pieces after the etching w re rins d with water and the surface states of thes test pieces w re examined by microanalysis with secondary ions but n impurity was detected.

Naturally, various embodim nts xplained in conn ction with 1 by making ref rence to Figs. 1A and 1B, Figs. 2A and 2B, Figs. 3A to 3C and Figs. 4A to 4C can be realized als in the cas where the mixture solution of hydrofluoric acid and aqu ous hydrog n peroxide is used as the etching solution.

## H(4)

A description will now be given of the case where a mixture of hydrofluoric acid, an alcohol and aqueous hydrogen peroxide is used as the electroless wet chemical etching solution for porous SI, with reference to Fig. 70.

Fig. 7D shows the time dependency of etched thickness of porous SI as observed when the porous SI is immersed in a mixture liquid of hydrofluoric acid, ethyl alcohol and aqueous hydrogen peroxide without aglitation of the liquid. The porous SI was formed by anodizing monocrystalline SI under the conditions shown below. The use of the monocrystalline SI as the starting material for forming the porous SI structure through anodization is only illustrative and SI of other crystalline structures can be used as the starting material.

Voltage applied: 2.6 (V)

Current density: 30 (mA·cm<sup>-2</sup>)
Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

Time: 2.4 (hours)
Thickness of porous Si: 300 (μm)

Porosity: 56 (%)

Test pieces of the porous Si prepared as described above were immersed, without agitation, in a mixture solution of 49 % hydrofluoric acid, ethyl alcohol and aqueous hydrogen peroxide (10: 6: 50) (white circles) and in a mixture solution of 49 % hydrofluoric acid, ethyl alcohol and hydrogen peroxide (10: 2: 10) (black circles). The reductions in the thicknesses of the porous SI test pieces were then measured. Large rates of atching of the porous Si were observed: namely, in the case of the 10: 6: 50 solution, the porous Si was etched by 107 µm and, in case of the 10: 2: 10 solution, the porous Si was etched by 120 µm, in about 40 minutes. After elapse of 80 minutes, the porous Si was etched by a thickness as large as 244 µm in the case of the 10: 5: 50 solution, with high degrees of states of the etched surfaces. The concentration of aqueous hydrogen peroxide was 30 % in this case but the hydrogen peroxide concentration may be determined in a range which provides an appreciable effect of addition of hydrogen peroxide and which does not cause any practical problem in the production process.

The etching rate has dependencies on the density of the hydrifluoric acid solution, as well as on the temperature of the same. The addition of alcohol serves to accelerate oxidation of silicon, thus enhancing the reaction speed as compared to the case where hydrogen peroxide is not used. It is also possible to control the reaction speed by suitably selecting the content of hydrogen peroxide. On the other hand, the addition of alcohol serves to remove, without delay, bubbles of reaction gases generated as a result of the etching from the etched surface, without requiring sgitation of the solution, thus offering high degrees of efficiency and uniformity of etching of the porous Si.

The solution density and the solution temperature are determined such that a practical etching speed is obtained and such that the effect of the use of hydrofluoric acid, alcohol and hydrogen peroxide is appreciable, although the mixture solutions of 49 % hydrofluoric acid, ethyl alcohol and hydrogen peroxide (10 : 6 : 50) and 49 % hydrofluoric acid, ethyl alcohol and hydrogen peroxide (10 : 2 : 10), as well as the room temperature as the solution temperature, are mentioned, these solution densities and temperature are only illustrative and are not intended to restrict the scope of the invention.

The HF concentration with respect to the etching solution preferably ranges between 1 and 85 %, more preferably between 5 and 80 % and most preferably between 5 and 80 %. The concentration of H<sub>2</sub>O<sub>2</sub> with respect to the etching solution is preferably between 10 and 95 %, more preferably between 10 and 80%, and is determined so as to provide an appreciable effect of addition of the hydrogen peroxide. The concentration of the alcohol with respect to the etching solution is preferably determined to be 80 % or less, more preferably 60 % or less and most preferably 40 % or less, and is selected so as to provide an appreciable effect of addition of the alcohol. The temperature is selected to range preferably 60 to 100°C, more preferably 50 80°C and most preferably 50 60°C.

Alcohol to be used in the present invention is not limited to ethyl alcohol and includes those alcohols such as isopr pyl alc hol which can practically be used in preparation proc as and accomplish the

The porous Si after the etching was rins d with water and the rinsed surface was xamined by microanalysis by using s condary ions but n Impurity was d tected.

This type of etching solution is advantageous in that bubbl is of reaction product gases gin rated as a result of thill etching can be removing delay from the surface being itching, dufficing agitation, so that this surface is etched with high digrees of smoothness and uniformity in the total surface.

A description will now be given of the etching characteristics of porous SI and non-porous SI as observed when they are etched by a mixture solution of hydrofluoric acid, ethyl alcohol and aqueous hydrogen peroxide, with specific reference to Fig. 6D.

Fig. 6D shows the time dependencies of etched thickness of porous SI and monocrystalline SI as observed when the porous SI and the monocrystalline SI are immersed in a mixture solution of hydrofluoric acid, ethyl alcohol and aqueous hydrogen peroxide, without aglation. The prorus SI was formed by anodization of monocrystalline SI conducted under the same conditions as those shown before. The use of monocrystalline SI as the starting material for forming porous SI throught anodization is only illustrative and SI of other crystalline structures can be used as the starting material.

A lest piece of porous SI prepared as described above was immersed, without agitation, in a mixture solution of 49 % hydrofluoric acid, eithy alcohal and aqueeus hydrogen peroxide (10 is : 50) (while icrides) at the norm temperature, and reduction in the thickness of the porous SI was measured. The porous SI was rapidly etched: namely, by a thickness of 107 µm in 40 minutes and 244 µm in 80 minutes, with high degrees of surface quality and uniformity. Although the concentration of the aqueeus hydrogen peroxide was 30 % in this case, the content of hydrogen peroxide may be suitably determined within a range which does not impair the effect of addition of hydrogen peroxide and which does not cause any practical inconvenience in the production.

The etching rate depends on the concentration and the temperature of the hydrofluoric acid and aqueous hydrogen peroxide.

The addition of hydrogen peroxide serves to accelerate oxidation of silicon, thus enhancing the reaction speed as compared to the case where hydrogen peroxide is not added. Furthermore, the reaction speed can be controlled by suitably selecting the content of the hydrogen peroxide. In addition, alcohol serves to remove, without delay, bubbles of the reaction product gases generated as a result of the etching without requiring agitation, thus ensurina high degrees of uniformity and etching of the procus of the procuss of

A test piece of a non-porous SI of 500 µm thick was immersed in a mixture solution of 49 % hydrofiluoric acid, ethyl alcohol and aqueous hydrogen peroxide (10 : 8 : 50) (black circles) at the room temperature, without agitation of the solution. The reduction in the thickness was then measured. In this case, the test piece of nonporous SI was etched only by 100 Angstrom or less even after elapse of 120 minutes. The etching rate showed almost no dependency on the solution concentration and temperature.

Both the porous and non-porous Si test pieces after the etching were rinsed with water and the surface states of these test pieces were examined by microanalysis with secondary lons but no impurity was detected.

Naturally, the various embodiments explained in connection with I by making reference to Figs. 1A and IB, Figs. 2A and 2B, Figs. 3A to 3C and Figs. 4A to 4C can be realized also in the case where the mixture solution of hydroffuoric acid, alcohol and aqueous hydrogen peroxide is used as the etchant for porous SI.

I-(5)

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A description will now be given of the case where a buffered hydrofluoric acid is used as the electroless wet chemical etching solution for porous Si, with reference to Fig. 7E. For instance, an aqueous solution containing 36.2 % of ammonium fluoride (NH<sub>4</sub>F) and 4.5 % of hydrogen fluoride (HF) is used as the buffered hydrofluoric acid.

Fig. 7E shows the time dependency of etching thickness of porous Si as observed when the porous Si as immersed in the buffered hydrofluoric acid followed by agitation. The porous Si was formed by anodizing monocrystalline Si under the conditions shown below. The use of the monocrystalline Si as the starting material for forming the porous Si structure through anodization is only illustrative and Si of other crystalline structures can be used as the starting material.

Voltage applied: 2.6 (V)

Current density: 30 (mA-cm<sup>-2</sup>)

Anodizing solution:  $HF:H_2O:C_2H_5OH = 1:1:1$ 

2.4 (hours)

5 Time:

Thickness of porous Si: 300 (um)

Porosity: 56 (%)

Test pic s of the porous Siprepared as discribed abovewire imm rs d, followed by agitation, in the

buffered hydrofluoric acid (white circles) and in a 20 % diluted buff red hydrofluoric acid (black circl s). The reductions in thi thickin sees of thi porous SI test pleaces where the measured. Large rates of etching of this porous SI were observed: namily, in the case of buff red hydrofluoric acid, the porous SI was etched by my man, in case of the 20 % diluted buffered hydrofluoric acid, the porous SI was to the dispersion of the dispersion of the case of the dispersion of the case of the buff red hydrofluoric acid and 94 µm in the case of the buff red hydrofluoric acid, with high degrees of states of the etched surfaces.

The etching rate has dependencies on the density of the hydrofluoric acid solution, as well as on the temperature of the same. The density of the solution and the temperature of the same are determined to fall within the ranges which would not cause any practical inconvenience. Although the buffered hydrofluoric acid which is an aqueous solution containing 36.2 % of ammonium fluoride (NH,F) and 4.5 % of hydrogen fluoride (HF) and the 20 % diluted buffered hydrofluoric acid, as well as the room temperature as the solution temperature, are mentioned, these solution densities and temperature are only illustrative and are not intended to restrict the scope of the invention.

The HF concentration in the buffered hydrofluoric acid with respect to the etching solution preferably ranges between 1 and 95 %, more preferably between 1 and 85 % and most preferably between 1 and 70 %. The concentration of NH<sub>4</sub> in the buffered hydrofluoric acid with respect to the etching solution is preferably selected to range between 1 and 95 %, more preferably between 5 and 80% and most preferably between 5 and 80%. The temperature is selected to range preferably 0 to 100°C, more preferably 5 to 80°C and most preferably 5 to 80°C and most preferably 5 to 80°C.

The porous Si after the etching was rinsed with water and the rinsed surface was examined by microanalysis by using secondary ions but no impurity was detected.

A description will now be given of the etching characteristics of porsus Si and non-porsus SI as observed when they are etched by the buffered hydrofluoric acid, with specific reference to Fig. 6E. Fig. 6E shows the time dependencies of etching of porsus SI and monocrystalline SI as observed when the porsus SI and the monocrystalline SI are immersed in the buffered hydrofluoric acid. The porsus SI was formed by anodization of monocrystalline SI are immersed in the buffered hydrofluoric acid. The porsus SI was formed by anodization of monocrystalline SI conducted under the same conditions as those shown before. The use of monocrystalline SI as the starting material is forming porsus SI through anodization is only illustrative and SI of other crystalline structures can be used as the starting material.

A test piece of porous SI prepared as described above was immersed, followed by agitation, in the buffered hydrofluoric acid (white circles) at the room temperature, and reduction in the thickness of the porous SI was measured. The porous SI was rapidly etched: namely, by a thickness of 70 µm in 40 minutes and 118 µm in 120 minutes, with high degrees of surface quality and uniformity.

The etching rate has dependencies on the density of the hydrofluoric acid solution, as well as on the temperature of the same. The density of the solution and the temperature of the same are determined to fall within the ranges which would not cause any practical inconvenience. Although the buffered hydrofluoric acid which is an aqueous solution containing 38.2 % of ammonium fluoride (NH<sub>2</sub>F) and 4.5 % of hydrogen fluoride (HF) as well as the room temperature, as the solution temperature, are mentioned, these solution densities and temperature are only illustrative and are not intended to restrict the scope of the invention.

A test piece of a non-porous Si of 500 µm thickness was immersed in the buffered hydrofluoric acid(black circles) at the room temperature, followed by agitation of the solution. The reduction in the thickness was then measured. In this case, the test piece of non-porous Si was etched only by 100 Angstroms or less even after slapse of 120 minutes. The etching rate showed almost no dependency on the solution density and tempera-

Both the porous and non-porous SI test pieces after the etching were rinsed with water and the surface states of these test pieces were examined by microanalysis with secondary ions but no impurity was detected.

Obviously, various etching methods explained in connection with (1) by making reference to Figs. 1A and 1B; 28. 2A and 28. Figs. 3A to 3C and Figs. 4A to 4C can be realized also in the case where the buffered hydrofluoric acid is used as the etching solution.

I-(6)

45

A description will now be given of the case where a mixture liquid of a buffered hydrofluoric acid and an alcohol is used as the electroless wet chemical atching solution for porous SI, with reference to Fig. 7F. For instance, an aqueous solution containing 36.2 % of ammonium fluoride (NH<sub>4</sub>F) and 4.5 % of hydrogen fluoride (HF) is used as the buffered hydrofluoric acid.

Fig. 7F shows the time dependency of etching thickness of porous Si as observed wh n the porous Si is immersed in the mixed solution of the buff red hydroflu ric acid and thyfalcoh I, without agitation. The porous

SI was f med by anodizing monocrystalline SI und r th conditions shown b low. The use of th monocrystalline SI as th starting material for forming the porcus SI structure through anodization is only flustrativ and SI of other crystalline structures can be used as the starting material.

Voltag appli d: 2.6 (V)

Curr nt d nsity: 30 (mA-cm<sup>-2</sup>)

Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

Time: 2.4 (hours)
Thickness of porous Si: 300 (µm)
Pomsity: 56 (%)

Porosity: 56 (%)
Test places of the porous Si prepared as described above were immersed, without agitation, in a mixture solution of the buffered hydrofluoric acid and ethyl alcohol(10:1) (white circles) and in a mixture solution of 20 % diluted buffered hydrofluoric acid and ethyl alcohol(10:1) (black circles). The reductions in the thicknesses of the porous Si test please were then measured. Large rates of etching of the porous Si were observed: namely, in the case of the mixture solution of the buffered hydrofluoric acid and ethyl alcohol (10:1), the porous Si was etched by 67 µm and, in case of the mixture solution of the 20 % diluteb buffered hydrofluoric acid and ethyl alcohol (10:1), the porous Si was etched by a thickness as large as 112 µm, in about 40 minutes. After elapse of 120 minutes, the porous Si was etched by a thickness as large as 112 µm in the case of the mixture solution of Dx 6 filluteb buffered hydrofluoric acid and ethyl alcohol (10:1), with high degrees of states of the etched surfaces. The etching rate has dependencies on the density of the hydrofluoric acid solution, as well as on the temperature of the same. The addition of alcohol serves to remove, without delay, bubbles of reaction product gases generated as a result of the etching from the surface being etched, without requiring agitation, thus enabling etching of the porous Si with high degrees of winformity and efficiency.

The density of the solution and the temperature of the same are determined to fall within the ranges which would not cause any practical inconvenience. Although the mixture solution of the buffered hydrofluoric acid and ethyl alcohol (10: 1) and the mixture solution of the 20 % diluted buffered hydrofluoric acid and ethyl alcohol (10: 1), as well as the room temperature as the solution temperature, are mentioned, these solution densities and temperature are only illustrative and are not intended to restrict the scope of the invention.

The HF concentration in the buffered hydrofluoric acid with respect to the atching solution preferably ranges between 1 and 9.5%, more preferably between 1 and 8.5% and most preferably between 1 and 7.0%. The concentration of NH<sub>4</sub> in the buffered hydrofluoric acid with respect to the etching solution is preferably selected to range between 1 and 9.5%, more preferably between 5 and 8.0% and most preferably between 5 and 8.0%. The concentration of the alcohol with respect to the atching solution is preferably 8.0% or less, more preferably 6.0% or less and most preferably 4.0% or less, more preferably 6.0% or less and most preferably 5.0% or less and mos

Although ethyl alcohol has been specifically mentioned, other alcohols such as isopropyl alcohol, which does not cause any inconvenience in the commercial production and which can provide an appreciable effect of addition of auch alcohol, may be used as the alcohol used in this type of etching solution.

The porous SI after the etching was rinsed with water and the rinsed surface was examined by microanalysis by using secondary ions but no impurity was detected.

in this etching solution, bubbles of reaction product gases generated as a result of the etching can be removed without delay and without requiring agitation of the solution, by virtue of the addition of the alcohol, so that the bottoms of minute recesses can be formed with high degrees of smoothness and uniformity.

A description will now be given of the etching characteristics of porous SI and non-porous SI as observed when they are etched by the mixture solution of the buffered hydrofluoric acid and the ethyl alcohol, with specific reference to Fig. 6F. Fig. 6F shows the time dependencies of etching thickness of porous SI and monocrystalline SI as sobserved when the porous SI and the monocrystalline SI are immersed in the mixture solution of the buffered hydrofluoric acid and ethyl alcohol. The porous SI was formed by anodization of monocrystalline SI on ducted under the same conditions as those shown before. The use of monocrystalline SI as the starting material for forming porous SI through anodization is only illustrative and SI of other crystalline structures can be used as the starting material.

A test piece of porous Si prepared as described above was immersed, without agitation, in the mixture solution of the buffered hydrofluoric acid and ethyl alcohol (10:1) (white circles) at the room temperature, and reduction in the thickness of the porous Si was measured. The porous Si was rapidly etched: namely, by a thickness of 67 µm in 40 minutes and 112 µm in 120 minutes, with high degrees of surface quality and uniformity.

At st piece of an n-porous SI of 500  $\mu$ m thickness was immers d in the mixture solution of the buff red hydrofluoric acid and thyl alcohol (10:1) (black circles) at the room temperature, without agitation of the sol-

ution. The r duction in the thickness of th non-porous SI was then measured. In this cas , the test pi ce of non-porous SI was etched inly by 100 Angstroms or less v in aft relaps if 120 minutes. The tching rate showed almost no dep indency on the clutton of insity and temp rature.

Both th p rous and non-porous Si test pieces after the tching were rins d with water and th surface states of thes test pieces w re examined by microanalysis with secondary ions but no impurity was detected.

Obviously, various tching methods explained in connection with (1) by making reference to Figs. 1A and 18, Figs. 2A and 2B, Figs. 3A to 3C and Figs. 4A to 4C can be realized also in the case where the mixture solution of the buffered hydrofusoric acid and alcohol is used as the etching solution.

I-(7)

A description will now be given of the case where a mixture solution of a buffered hydrofluoric acid and hydrogen peroxide is used as the electroless wat chemical etching solution for porous SI, with reference to Fig. 7G For instance, an aqueous solution containing 36.2 % of ammonium fluoride (NH<sub>4</sub>F) and 4.5 % of hydrogen fluoride (NH<sub>4</sub>F) is used as the buffered hydrofluoric acid.

Fig. 7G shows the time dependency of etching thickness of porous SI as observed when the porous SI is missered in the mixed solution of the buffered hydrofluoric acid and hydrogen peroxide followed by agitation. The porous SI was formed by anodizing monocrystalline SI under the conditions shown below. The use of the monocrystalline SI as the starting material for forming the porous SI structure through anodization is only illustrative and SI of other crystalline structures can be used as the starting material.

Voltage applied: 2.6 (V)

Current density: 30 (mA-cm-2)

Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

Time: 2.4 (hours)
Thickness of porous Si: 300 (µm)

Thickness of porous Si: 300 (μm)
Porosity: 56 (%)

Test pieces of the porous SI prepared as described above were immersed, followed by agitation, in a mixture solution of the buffered hydrofluoric acid and hydrogen peroxide (1:5) (white circles) and in a mixture solution of the buffered hydrofluoric acid and hydrogen peroxide (5:1) (black circles). The reductions in the thicknesses of the porous SI test pieces were then measured. Large rates of stching of the porous SI west observed: namely, in the case of the 1:5 mixture solution, the porous SI was etched by 8 µm and, in case of the 5:1 mixture solution, the porous SI was etched by 8 µm and, in case of the 5:1 mixture solution, with high degrees of states of the etched by a mixture solution, with high degrees of states of the etched surfaces. In this case, the concentration of hydrogen peroxide was 30 %. This, however, is only illustrative and the concentration of hydrogen peroxide was 30 %. This, however, is only illustrative and the concentration of hydrogen peroxide. The etching rate has dependencies on the solution densities of the buffered hydrofluoric acid and hydrogen peroxide, as well as on the temperature of the same. The addition of hydrogen peroxide accelerates the oxidation of sillicon, thus attaining a higher reaction speed as compared to the case where hydrogen peroxide is not added. In addition, the reaction speed can be controlled by suitably determining the content of hydrogen peroxide.

The density of the solution and the temperature of the same are determined to fall within the ranges which would not cause any practical inconvenience in commercial production. Although the mixture solution of the buffered hydrofluoric acid and hydrogen peroxide (1:5) and the mixture solution of the buffered hydrofluoric acid and hydrogen peroxide (5:1), as well as the room temperature as the solution temperature, are mentioned, these solution densities and temperature are only illustrative and are not intended to restrict the scope of the invention.

The HF concentration in the buffered hydrofluoric acid with respect to the etching solution preferably ranges between 1 and 95 %, more preferably between 1 and 85 % and most preferably between 1 and 70 %. The concentration of NN<sub>L</sub> in the buffered hydrofluoric acid with respect to the etching solution is preferably selected to range between 1 and 95 %, more preferably between 5 and 90% and most preferably between 5 and 80%. The concentration of  $\frac{1}{2}O_2$  with respect to the etching solution is preferably 10 95 %, more preferably 5 to 90% and most preferably 10 to 80 %, and is determined to make the effect of addition of the hydrogen peroxide. The temperature is selected to range preferably 0 to 100°C, more preferably 5 to 80°C and most preferably 5 to 80°C and most preferably 5 to 80°C and most preferably 5 to 80°C.

The porous SI after the etching was rinsed with water and the rinsed surface was examined by microanalysis by using secondary ions but no impurity was direct id.

A d scription will now b giv n of th tching characteristics f porous Si and non-porous Si as obs rv d

wh n they ar etch d by the mixture solution of the buffered hydrofluoric acid and hydrogen peroxid , with specific refer note to Fig. 6G. Fig. 6G shows the time of pendencies of etching thickness of porous SI and monocrystalline SI are immersed in the mixture soluti n of the buff red hydrofluoric acid hydrogen peroxid . The porous SI was formed by anodization of monocrystalline SI conducted und r th same conditi ns as those shown b forc. Thus of m nocrystalline SI conducted und r th same conditi ns as those shown b forc. Thus of m nocrystalline SI are the same condition of the same conditions are solved to the same should be sufficiently and SI of other crystallin structures can be used as the starting material.

A test plece of porous SI prepared as described above was immersed, followed by agitation, in the mbsture solution of the buffered hydrofluoric acid and hydrogen peroxide (1:5) (while circles) at the room temperature, and reduction in the thickness of the porous SI was measured. The porous SI was rapidly etched: namely, by a thickness of 88 µm in 40 minutes and 147 µm in 120 minutes, with high degrees of surface quality and uniformity. Although in this case the concentration of hydrogen peroxide was 30 %, this is only illustrative and the content of hydrogen peroxide is suitably selected within a range which does not cause any practical inconventience and which does not impair the effect produced by the addition of hydrogen peroxide.

Both the porous and non-porous Si test pieces after the etching were rinsed with water and the surface states of these test pieces were examined by microanalysis with secondary ions but no impurity was detected.

Obviously, various etching methods explained in connection with (1) by making reference to Figs. 1A and 18, Figs. 2A and 2B, Figs. 3A to 3C and Figs. 4A o4 Cc an be realized also in the case where the mixture solution of the buffered hydrolluoric acid and alcohol is used as the etching solution.

I-(8)

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The following will now be given of the case where a mixture solution of a buffered hydrofluoric acid, an alcohol and hydrogen peroxide is used as the electroless wet chemical etching solution for porous Si, with reference to Fig. 7H. For instance, an aqueous solution containing 38.2 % of ammonium fluoride (NH<sub>4</sub>F) and 4.5 % of hydrogen fluoride (HF) is used as the buffered hydrofluoric acid.

Fig. 7H shows the time dependency of etching thickness of porous SI as observed when the porous SI is immersed in the mixed solution of the buffered hydrofluoric acid, ethyl alcohol and hydrogen peroxide without agitation. The porous SI was formed by anotizing monocrystalline SI under the conditions shown below. The use of the monocrystalline SI as the starting material for forming the porous SI structure through anodization is only illustrative and SI of other crystalline structures can be used as the starting material to the starting material starting material.

Voltage applied: 2.6 (V)

Current density: 30 (mA·cm<sup>-2</sup>)

Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

Time: 2.4 (hours)
Thickness of porous Si; 300 (μm)

Porosity: 56 (%)

Test pieces of the porous Si prepared as described above were immersed, without agitation, in a mixture solution of the buffered hydrofluoric acid, ethyl alcohol and hydrogen peroxide (10:6:50) (white circles) and in a mixture solution of the buffered hydrofluoric acid, ethyl alcohol and hydrogen peroxide (50:6:10) (black circles). The reductions in the thicknesses of the porous SI test pieces were then measured. Large rates of etching of the porous SI were observed: namely, in the case of the 10:6:50 mixture solution, the porous SI was etched by 83 µm and, in case of the 50; 6; 10 mixture solution, the porous Si was etched by 100 µm, in about 40 minutes. After elapse of 120 minutes, the porous Si was etched by a thickness as large as 140 µm in the case of the 10:6:50 mixture and 168 µm in the case of the 50:6:10 mixture solution, with high degrees of states of the etched surfaces. In this case, the concentration of hydrogen peroxide was 30 %. This, however, is only illustrative and the concentration of hydrogen peroxide is suitably selected within a range which does not impair the effect of addition of hydrogen peroxide. The etching rate has dependencies on the concentrations of the buffered hydrofluoric acid and hydrogen peroxide, as well as on the temperature of the same. The addition of hydrogen peroxide accelerates the oxidation of silicon, thus attaining a higher reaction speed as compared to the case where hydrogen peroxide is not added. In addition, the reaction speed can be controlled by suitably determining the ratio of hydrogen peroxide. The addition of alcohol enables bubbles of reaction products gases generated as a result of the etching to be removed from the surface being etched, without delay and without aditation, thus making it possible to etch the porous SI uniformly and with high efficiency.

The concentrations of the solution and the temperature of the solution are determined to fall within the ranges which provide the above effects of the use of the buffered hydrofluoric acid, hydrogen peroxide and the alcohol and which would not cause any practical inconvenience in commercial production. Although the mixture soluti no f the buffered hydrofluoric acid, ethyl alcohol and hydrogen peroxid (10:8:50) and th mixture solution.

ution fith buff red hydrofluoric acid, thyl alcohol and hydrog in peroxide (50:6:10), as will as the room temperature as the soution it imperature, are mention in these solution ratio and temperature are only illustrative and are not intend if to restrict this scope of this invention.

Th HF concentration in th buffered hydrofluoric acid with respect to th tching solution preferably ranges between 1 and 95 %, more prefer rably by tween 1 and 85 % and most pref rably between 1 and 70 %. The concentration of NH<sub>2</sub>Fin th buffered hydrofluoric acid with respect to the citching is lution is pref rably selected to range between 1 and 95 %, more preferably between 5 and 90 % and most preferably between 5 and 90 % and most preferably between 5 and 90 % and most preferably 10 to 80 %, and is determined to make the effect of addition of the alcohol appreciable. The concentration of the alcohol with respect to the etching solution is preferably 80 % or less, more preferably 60 % or less and most preferably 40 % or less, and is determined to make the effect of addition of the alcohol appreciable. The temperature is selected to range preferably 0 to 100°C, more preferably 5 to 80°C and most preferably 5 to 80°C and most preferably 5 to 80°C.

Although ethyl alcohol has been specifically mentioned, other alcohols such as isopropyl alcohol, which does not cause any inconvenience in the commercial production and which can provide an appreciable effect of addition of such alcohol, may be used as the alcohol used in this type of etching solution.

The porous SI after the etching was rinsed with water and the rinsed surface was examined by microanalysis by using secondary ions but no impurity was detected.

This etching solution enables bubbles of reaction product gases generated by the etching to be removed from the surface being etched, without delay and without requiring agitation, so that the etching can be performed with high degrees of smoothness and uniformity at the bottoms of minute recesses of the surface to be etched.

Clearly, embodiments explained in the above (1) by making reference to Figs. 1A and 1B, Figs. 2A and 2B, Figs. 3A to 3C and Figs. 4A to 4C can be realized also in the case where the mixture solution of the buffered hydrofluoric acid, alcohol and hydrogen peroxide is used as the etching solution.

II. The following will now be given of a process of the invention for producing a semiconductor member.

As explained before, the first embodiment of the process for producing the semiconductor member in accordance with the present invention has the following features.

Namely, the first embodiment of the process of the invention for producing a semiconductor member comprises the steps of forming a member having a non-porous silicon monocrystalline layer and a porous silicon layer, bonding to the monocrystalline layer a member having an insulating material surface; and removing by etching the porous silicon layer by immersing it in hydrofluoric add.

As explained before, the second embodiment of the process of the invention for producing a semiconductor member uses the same steps as those in the method of the first embodiment, except that, in place of the hydrofluoric acid used in the first embodiment, one of the second to eighth embodiment: of the etching solutions mentioned before.

The third to sixth embodiments of the process of the Invention for producing a semiconductor member, which also were explained before, are more practical embodiments of the first and the second embodiments of the process of the Invention. The process of the present invention for producing a semiconductor member will be described hereinafter with reference to the third to sixth embodiments.

## ii-(1)

The third embodiment of the process of the invention for producing a semiconductor member will be described with reference to the drawings.

# Embodiment 1

The following will be first given of a method in which the whole member is changed into porous structure and then a monocrystalline layer is formed on the procus structure by epitaxial growth method. Figs. 8A to 8C are schematic sectional views of the semiconductor member illustrating each of steps of the process.

Referring to Fig. 8A, as the first step, an SI monocrystalline semiconductor member 11 is prepared and is wholly changed into porous structure and, then, an epitaxial growth method is applied to one surface of the porous member, thereby forming a thin film of monocrystalline SI layer 12. The porous structure of SI member is formed by, for example, an anodization employing an HF soluti n. The Initial monocrystallin SI having the density of 2.33 /gcm² can b changed into a por us SI member it it on they of which can be varied within th

range betw en 1.1 and 0.6 g/cm³ by varying the HF concentration of the etching solution betwoen 50 % and 20 %.

Referring now to Fig. 8B, a light-transmissiv substrat 13, which is typically a glass sheet, is prepared and bond d on th surface of the monocrystalline Sllayer 12 nt by prorus Si memb r. Subsequently, an Si<sub>3</sub>N<sub>4</sub> lay r14 is formed by d position as an anti-teching film to cov r th mitter member composed of the lay r12 and the substrat 13 and the Si<sub>3</sub>N<sub>4</sub> layer on th porous Si memb r11 is removed. Although Si<sub>3</sub>N<sub>4</sub> layer is sultably used as the anti-etching layer. It is possible to use other materials such as Aplezon wax as the materials of the anti-etching layer. The porous Si member 11 is then immersed in the etching solution of the present invention and the solution is agitated so that only the porous Si is etched by electricless chemical etching, whereby a thinned non-procus monocrystalline sillicon layer 12 is left on the light-transmissive substrate 13.

Fig. 8C shows the semiconductor member obtained by the present process, it will be sald that, as a result of the removal of the anti-othing Si,N, isyer 14 in the step shown in Fig. 8B, a monocrystalline SI layer 12 having a crystallinity equivalent to that of a silicon wafer is formed on the light-transmissive substrate 13 with high degrees of smoothness and uniformity and with a small thickness, over a wide area covering the whole surface of the wafer.

The semiconductor member thus obtained is advantageous from the view point of production of an insulation-isolated electronic device.

## Embodiment 2

The following will now be given of a process in which an N-type layer is formed before changing the Initial member into porous structure and, subsequently to the formation of the P-type layer, a selective anodization is effected to change only the P-type substrate into prorus structure.

Referring to Fig. 9A, as the first step, a layer 32 of a low impurity concentration is formed on the surface of a P-type Si monocrystalline substrate 31, by an epitaxial growth. Alternatively, an N-type monocrystalline layer 32 may be formed on the surface of the P-type Si monocrystalline substrate 31 by ion-implantation of pro-

Then, as shown in Fig. 9B, the P-type Si monocrystalline substrate 31 is changed into a porous Si substrate of the P-type Si monocrystalline substrate 31, an anodization using, for example, an HF solution. The initial monocrystalline SI having the density of 2.33 g/cm² can be changed into a porous member the density of which can be varied within the range between 1.1 and 0.6 g/cm² by varying the HF concentration of the etching solution between 50 % and 20 %.

Referring now to Fig. 9C, a light-transmissive substrate 34, which is typically a glass sheet, is prepared and bonded on the surface of the monocrystalline Si layer 32 on the porous Si member. Subsequently, an Si<sub>2</sub>N<sub>4</sub> layer 35 is formed by deposition as an anti-teching film to over the entire member composed of the layer 32 and the substrate 34 and the Si<sub>2</sub>N<sub>4</sub> layer on the porous Si member 33 is removed. Although Si<sub>2</sub>N<sub>4</sub> layer is subsubly used as the anti-teching layer, it is possible to use other materials such as Aplezon wax as the material of the anti-teching layer. The porous Si substrate 33 is then immersed in the etching solution of the present invention and the solution is aglitated so that only the porous Si is etched by electroless chemical etching, whereby a thinned non-porous monocrystalline silicon layer 32 is left on the light-transmissive substrate 34.

Fig. 9D shows the semiconductor member obtained by the present process. That is, as a result of the removal of the anti-stching SlyN, layer in the step shown in Fig. 9D, a monocrystalline SI layer 32 having a crystallinity equivalent to that of a silicon wafer is formed on the light-transmissive substrate 34 with high degrees of smoothness and uniformity and with a small thickness, over a wide area covering the whole surface of the

The semiconductor member thus obtained is advantageous from the view point of production of an Insulation-isolated electronic device.

According to the result of an observation by a transmission electron microscope, micro-pores of an average distinct of about 800 Angstrom are formed in the porous SI layer, so that the density of the layer has been reduced half or below that of the monocrystalline SI.

Nevertheless, the monocrystallinity is still maintained, so that it is possible to form a monocrystalline SI layer on the porous layer by epitaxial growth. When the temperature exceeds 100°C, rearrangement of the pres occurs, which impedes the acceleration of the etching. For this reason, the epitaxial growth of the SI layer is preferably effected by a low-temperature growth method such as, for example, a molecule-ray epitaxial growth method, a CVD method such as plasma CVD method, low-pressure CVD method or photo-CVD method, a bias souther method or a liquid-phase growth method.

# 11-(2)

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The fourth imbodiment of the process of the lnv intion of producing a simiconductor member will be discribed with reference to the drawings.

#### Embodim nt 1

The following will be first given of a form in which the whole P- or high-density N-type substrate is changed into porous structure and then a monocrystalline layer is formed on the porous structure by epitaxial growth method Figs. 10A to 10C are schematic sectional views of the semiconductor member illustrating each of steps of the process.

Referring to Fig. 10A, as the first step, an SI monocrystalline semiconductor member 11 of P-type (or highdensity N-type) is prepared and is wholly changed into porous structure and, then, an epitaxial growth is effected by a suitable method on the surface of the porous member, thereby forming a thin film of monocrystalline SI layer 12. The porous structure is formed by, for example, an anodization employing an HF solution. The limital monocrystalline SI having the density of 2.33 g/cm² can be changed into a porous member the density of which can be varied within the range between 1.1 and 0.6 g/cm² by varying the HF concentration of the etching solution between 50 % and 20 %.

Referring now to Fig. 10B, another SI substrate 13 is prepared and an insulating layer (silicon oxide layer) 14 formed on the surface of this SI substrate 13. Subsequently, the surface of the insulating layer 14 of the SI substrate 13 is bonded to the surface of the monocrystalline layer 12 on the prorus SI substrate. The whole structure 11-14 composed of the substrates and layers to 14 is then immersed in the etching solution of the present invention and the solution is agitated so that only the porous SI is etched by electroless wet chemical etching, whereby a thinned non-porous monocrystalline silicon layer 12 is left on the insulating layer 14.

Fig. 10C shows the semiconductor member obtained by the present process. That is, the monocrystalline SI layer 12 having a crystallinity equivalent to that of a silkon wafer is formed on the insulating layer 14 on the SI substrate 13 with high degrees of smoothness and uniformity and with a small thickness, over a wide area covering the whole surface of the wafer.

The semiconductor member thus obtained is advantageous from the view point of production of an insulation-isolated electronic device.

#### Embodiment 2

The following will now be given of a process in which an N-type layer is formed before changing the initial member into porous structure and, subsequently to the formation of the N-type layer, a selective anodization is effected to change only the P-type substrate or the high-density N-type substrate into porous structure. Figs. 114 to 110 show, in schematic sectional views, the semiconductor member in different steps of the production process.

Referring to Fig. 11A, as the first step, a layer 22 of a low impurity concentration is formed on the surface of a P-type (or high-density N-type) Si monocrystalline substrate 21, by an epitaxial growth performed by a sultable method. Alternatively, an N-type monocrystalline layer 22 may be formed on the surface of the P-type Si monocrystalline substrate 21 by ion-implantation of proton.

Then, as shown In Fig. 118, the P-type Si monocrystalline substrate 21 is changed into a porous SI substrate 23 by effecting, on the reverse side of the P-type monocrystalline substrate 21, an anodization using, for example, an HF solution. The initial monocrystalline SI having the density of 2.33 g/cm³ can be changed into a porous member the density of which can be varied within the range between 1.1 and 0.6 g/cm³ by varying the HF concentration of the etching solution between 50 % and 20 %.

Referring now to Fig. 11C, another Si substrate 24 is prepared and an insulating layer 25 (eillicon oxide layer) is formed on the surface of the Si substrate 24. Then, the insulating layer 25 on the Si substrate 24 is bonded to the surface of the monocrystalline Si layer 22 on the porous substrate. Subsequently, the whole structure composed of the substrates and layers 22 to 25 is immersed in the etching solution of the present invention and the solution is agalted so that only the porous Si is etched by electroless chemical etching, whereby a thinned non-porous monocrystalline sillicon layer 22 is left on the insulating layer 25.

Fig. 11D shows the semiconductor member obtained by the present process. That is, a monocrystalline Si layer 22 having a crystallinity equivalent to that of a silicon wafer is formed on the insulating layer 25 with high degrees of smoothness and uniformity and with a small thickness, over a wide area covering the whole surface f the wafer.

The semiconductor member thus obtained is advantageous from the view point of production of an insu-

lation-is lated electronic d vic .

# 11-(3)

The fifth embodiment of th process of th Invention will b described with reference to th drawings. A described will be first given of a form in which th wholt SI substat is changed into porous structure and then a monocystalline layer is formed on the porous structure by epitaxial growth method.

Referring to Fig. 12A, as the first step, an Si monocrystalline substrate 11 is prepared and is wholly changed into procus structure and, then, an epitaxial growth is effected by a suitable method on the surface of the porous substrate, thereby forming a thin film monocrystalline layer 12. The porous structure is formed by, for example, an anodization employing an HF solution. The initial monocrystalline Si having the density of 2.33 g/cm² can be changed into a porous Si layer the density of which can be varied within the range between 1.1 and 0.6 g/cm² by varying the HF concentration of the etching solution between 50 % and 20 %. The porous layer is tended to form in a P type SI substrate. A transmission electromicroscopic observation showed that the porous SI layer thus formed has microproses of a mean diameter of about 600 Angstroms.

Referring now to Fig. 12B, a light-transmissive substrate 13, which is typically a glass sheet, is prepared. Then, the surface of the monocrystalline SI layer on the porous SI substrate is oxidized to form an oxide layer 14. The above-mentioned light-transmissive substrate 13 is then bonded on the surface of the oxide layer 14. This oxide layer plays an important role formation of device. Namely, with such an oxide layer, the interface level generated at the interface under the SI active layer can be made lower as compared with the glass interface, so that the characteristics of the electronic device can be remarkably improved.

Referring further to Fig. 12B, and Si<sub>2</sub>N<sub>4</sub> layer 15 is deposited as an anti-etching film(protective material) to cover the entire member composed of the two substrates bonded together, and the Si<sub>2</sub>N<sub>4</sub> layer on the surface of porous Si substrate is removed. Although Si<sub>2</sub>N<sub>4</sub> layer is suitably used as the anti-etching layer, it is possible to use other materials such as Apiezon wax as the material of the anti-etching layer. The porous SI substrat 1s then immersed in the etching solution of the present invention with agilating so that only the provost SI is etched by electroless chemical etching, whereby a thinned monocrystalline silicon layer is left on the light-transmissive substrate 13.

Fig. 12C shows the semiconductor member obtained by the described process. It will be seen that, as a result of the removal of the anti-etching Si<sub>2</sub>N<sub>4</sub> layer 15 in the step shown in Fig. 12B, a monocrystalline Si layer 12 having a crystallinity equivalent to that of a silicon wafer is formed on the light-transmissive substrate 13 with high degree of smoothness and uniformity and with a small thickness, over a wide area covering the whole surface of the wafer.

The semiconductor member thus obtained is advantageous from the view point of production of an insulation-isolated electronic device.

## 11-(4)

The sixth embodiment of the process of the invention for producing a semiconductor member will now be

## Embodiment 1

A description will be first given of a form in which the whole Si substrate is changed into porous structure and then a monocrystalline layer is formed by epitaxial growth method.

Figs. 13A to 13C illustrate successive steps of the first embodiment in accordance with the invention. Referring to Fig. 13A, as the first step, an 5 imonocystalline substrate is prepared and is wholly changed into porous structure (11). Then, an epitaxial growth is effected by a suitable method on the surface of the porous substrate, thereby forming a thin film of monocystalline layer 12. The porous structure is formed by, for example, an anodization employing an HF solution. The initial monocystalline SI having the density of 2.33 g/cm<sup>2</sup> can be changed into a porous SI layer the density of which can be varied within the range between 1.1 and 0.6 g/cm<sup>2</sup> by varying the HF concentration of the etching solution between 50 % and 20 %. A transmission electromicroscopic observation showed that the porous SI layer thus formed has micropores of a mean diameter of about 600 Anostroms.

Referring now to Fig. 13B, another SI substrate 13 is prepared and an insulating material 14 is formed on the surface. Then the SI substrate having the insulating material 14 is bonded to the surface of an oxide layer 15 which is formed on th monocrystalline SI layer carri d by the por us SI substrate. The insulating material 14 may be a deposited silicon oxide, nitride, nitrided oxide, r tantalum, n t to mention the insulating layer of

Si. Th bonding step may b conducted by adh ring closely the rinsed surfac s, and heating both substrate in an oxyg n atmosph reo r a nitrog n atmosph re. Th oxid layer 15 is formed for the purpose of reducing th Interface level of the monocrystallin layer 12 which is the final active layer.

Th n, as shown in Fig. 13C, th porcus SI substrate 11 is immersed in the tching solution of the press invention and th solution is egistated, so that nly the porcus SI is tched by lectroless wet ch mical tribing so as to 1 av a thinned monocrystallin SI lay r on the insulating mat rial. Fig. 13C shows the semiconductor substrate obtained according to the present invention. As a result, a monocrystalline SI layer 12 having crystallinity equivalent to that of a silicon water is formed on the insulated substrate 13 through the intermediary of the insulating material 14 and the oxide layer 15, with high degrees of smoothness and uniformity and with a small thickness over a wide area covering the whole surface of the wafer.

The semiconductor member thus obtained is advantageous from the view point of production of an insuiation-isolated electronic device.

#### Embodiment 2

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A second embodiment will be described with reference to the drawings.

Figs. 14A to 14D show, in schematic sectional views, the second embodiment according to the present invention.

Referring to Fig. 14A, as the first step, a layer 32 of a low impurity concentration is formed on the surface of a P-type SI monocrystalline substrate 31, by an epitaxial growth performed by a suitable method. Alternatively, an N-type monocrystalline layer 32 may be formed on the surface of the P-type SI monocrystalline substrate 21 by implantation of proton.

Then, as shown in Fig. 14B, the P-type Si monocrystalline substrate 31 is changed into a porous Si substrate 33 by effecting, on the reverse side of the P-type Si monocrystalline substrate 31 by anotization using, for example, an HF solution. The initial monocrystalline Si having the density of 2.33 g/cm² can be changed into a porous member the density of which can be varied within the range between 1.1 and 0.6 g/cm² by varying the HF concentration of the etching solution between 50 % and 20 %. As explained before, this porous layer is formed in the P-type substrate.

Referring now to Fig. 14C, another Si substrate 34 is prepared and an insulating layer 35 is formed on the surface of the Si substrate 34. Then, the Si substrate 34 having the insulating layer 35 is bonded to the surface of the oxide layer 36 formed on the monocytabiline Si layer on the porous Si substrate. Then, the porous Si substrate is immersed in the etching solution of the present invention and the solution is agitated so that only the porous Si is atched by electroless chemical etching, whereby a thinned non-porous monocrystalline silicon layer is left on the insulation layer.

Fig. 14D shows the semiconductor substrate obtained by the described process. It will be seen that a monocrystalline SI layer 32 having a crystallinity equivalent to that of a silicon wafer is formed on the insulated substrate 34 through the intermediary of the oxide layer 36 and the insulation layer 35, with high degrees of smoothness and uniformity and with a small thickness, over a wide area covering the whole surface of the wafer.

The semiconductor member thus obtained is advantageous from the view point of production of an insulation-isolated electronic device.

The processes described above are the type in which the N-type layer is formed prior to changing into porous structure and then only the P-type substrate is selectively changed into porous structure by anodization.

III. The present invention will be described below in detail by way of examples. However, the invention is not limited to these examples except as defined in the appended claims.

## Example 1

A porous SI layer 21 was formed to a thickness of 50  $\mu$ m ( $t_2 = 50 \mu$ m) on the entirety of one of the major surfaces of a monocrystalline SI substrate 22 by anodization (Fig. 1A).

Anodization was performed under the following conditions:

Applied voltage: 2.6 (V)

Current density: 30 (mA·cm<sup>-2</sup>)
Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

Time: 0.4 (hour)

Thickness of porous Si: 50 (µm)
Porosity: 56 (%)

Thir aftir thip rous Si/monocrystallin. Si substrat, was subjected to selective, tching using a 49% HF

solution. In thirty-thre minutes, the porous SI was a lectively tched with the monocrystalline SI acting as an etch stopp r, only the monocrystalline SI being I ft b hind, as hown in Fig. 1B.

#### Exampi 2

Prior t anodization, boron ions were implanted in ne of th surfaces of a monocrystallin SI substrate 32 at an average concentration of  $1.0 \times 10^{10}$  cm<sup>-3</sup> in stripes spaced apart from each other by a distance of 100 ym. As shown in Fig. 2A, porous SI 31 was formed by anodization in stripes spaced apart from each other by a distance ( $b_3 = 100 \, \mu m$ ) of 100  $\mu m$ , each stripe having a width ( $a_5 = 100 \, \mu m$ ) of 100  $\mu m$  and a thickness ( $b_5 = 100 \, \mu m$ ) of 100  $\mu m$  and 100  $\mu$ 

Anodization was performed under the following conditions:

Applied voltage: 2.6 (V)

Current density: 30 (mA-cm<sup>-2</sup>)

Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

Thickness of porous Si: 1 (μm)

Porosity: 56 (%)

Thereafter, the porous St/monocrystalline Si substrate was subjected to selective etching using a 49% HF solution. In two minutes, the porous Si was selectively etched, only the monocrystalline Si being left behind, as shown in Fig. 2B.

#### Example 3

A 3 $\mu$ m ( $u_4$  = 3  $\mu$ m) thick polycrystalline Si layer 41 was formed on a monocrystalline SI substrate 42 by CVD (Fig. 3A). As shown in Fig. 3B, a surface layer of 2  $\mu$ m ( $t_4$  = 2  $\mu$ m) of the polycrystalline SI layer 41 was made porous by anodization to form a porous SIlayer 43.

Anodization was performed under the following conditions:

Applied voltage: 2.6 (V)

Current density: 30 (mA·cm<sup>-2</sup>)

Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>6</sub>OH = 1:1:1

Thickness of porous Si: 2 (μm)

Porosity: 56 (%)

Thereafter, the porous Si/polycrystalline Si/monocrystalline Si substrate was subjected to selective etching using a 49% HF solution. In four minutes, the porous Si was selectively etched with the polycrystalline Si selection as an etch shopper, only the polycrystalline Si and monocrystalline Si being left behind, as shown in Fig. 3C.

## Example 4

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A 3 $\mu$ m (u<sub>5</sub> = 3  $\mu$ m) thick polycrystalline Si layer 51 was formed on a monocrystalline SI substrate 52 by CVP. Prior to anodization, boron lons were implanted into the surface of the polycrystalline SI layer 51 at 1.0  $\times$  10 $^{10}$  cm<sup>-3</sup> in stripes spaced apart from each other by a distance of 20  $\mu$ m. As shown in Fig. 4A, porous SI 53 was formed by anodization in stripes spaced apart from each other by a distance (b<sub>5</sub> = 20  $\mu$ m) of 20  $\mu$ m, each stripe having a width (e<sub>8</sub> = 20  $\mu$ m) of 20  $\mu$ m and a thickness (g = 1  $\mu$ m) of 1  $\mu$ m.

Anodization was performed under the following conditions:

Applied voltage: 2.6 (V)

Current density: 30 (mA-cm<sup>-2</sup>)

Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>6</sub>OH = 1:1:1

Thickness of porous Si: 1 (µm)

Porosity: 56 (%)

Thereafter, the porous Sl/polycrystalline Sl/monocrystalline SI substrate was subjected to selective etching so using a 49% Hs odution. In two minutes, the porous SI was selectively etched, only the polycrystalline SI and monocrystalline SI being left behind, as shown in Fig. 4B.

## Example 5

A porous Si layer 61 was formed to a thickness of 50 µm (t<sub>6</sub> = 50 µm) on the entirety of one of the major surfaces of a monocrystalline SI substrate 62 by anodization (Fig. 5A).

Anodization was p rform d und r th following conditi ns:

Appli d voltage: 2.6 (V)

Current density: 30 (mA-cm<sup>-2</sup>)

Anodizing solution:  $HF:H_2O:C_2H_5OH = 1:1:1$ 

Time: 0.4 (hour)
Thickn ss of porous Si: 50 (µm)
Porosity: 56 (%)

As shown in Fig. 5B, a resist 63 was patterned in stripes spaced apart from each other by a distance ( $b_0$  = 100  $\mu$ m) of 100  $\mu$ m, each stripe having a width ( $a_0$  = 100  $\mu$ m) of 100  $\mu$ m.

Thereafter, the porous St/monocrystalline SI substrate was subjected to selective etching using a 49% HF solution. In thirty-three minutes, the produs SI was selectively removed, only the monocrystalline SI being left behind, as shown in Fig. 5C. Finally, the resist was removed (Fig. 5D).

## Example 6

Etching was performed in the same manner as that of Example 1 with the exception that a mixture solution (10:1) of 49% hydroffuoric acid and ethyl alcohol was used as an etchant. In twenty-nine minutes after initialization of etching, the prous SI was selectively removed with the monocrystatiline SI acid na san etch stopper.

## Example 7

Etching was performed in the same manner as that of Example 2 with the exception that a mixture solution (10:1) of 49% hydrofluoric acid and ethyl alcohol was used as an etchant. In one point seven minutes after initialization of etching, the porous SI was selectively removed, only the monocrystalline SI being left behind.

## Example 8

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Etching was performed in the same manner as that of Example 3 with the exception that a mixture solution (10:1) of 49% hydrofluoric acid and ethyl alcohol was used as an etchant. In three point four minutes after initialization of etching, the porous 5 was selectively removed with the polycrystalline SI acting as an etch stopper, only the polycrystalline SI and the monocrystalline SI being left behind, as shown in Fig. 3C.

#### Example 9

Etching was performed in the same manner as that of Example 4 with the exception that a mixture solution (10:1) of 49% hydroffuncir acid and ethyl alcohol was used as an etchant. In one point seven minutes after mitialization of etching, the porous Si was selectively removed, only the polycrystalline Si and monocrystalline Si being left behind, as shown in Fig. 4B.

#### Example 10

Etching was performed in the same manner as that of Example 5 with the exception that a mixture solution (10:1) of 49% hydroffuoric acid and ethyl alcohol was used as an etchant. In twenty-nine minutes after initialization of etching, the porous SI was selectively removed, only the monocrystalline SI being left behind, as shown in Fig. 5C. Finally. the resist was removed (Fig. 5D).

## 5 Example 11

Etching was performed in the same manner as that of Example 1 with the exception that a mixture solution (1:5) of 49% hydrofluoric acid and hydrogen peroxide was used as an etchant. In twenty-one minutes after initialization of etching, the porous SI was selectively removed with the monocrystalline SI acting as an etch stoper, only the monocrystalline SI being left behind, as shown in Fig. 18.

## Example 12

Etching was performed in the same manner as that of Example 2 with the exception that a mixture solution (1:5) of 49% hydrofluoric acid and hydrogen peroxide was used as an etchant. In one point three minutes after initialization of etching, the porous Si was selectively removed, only the monocrystalline Si being left behind, as shown in Fig. 2B.

## Exampl 13

Elching was p fromed in the sam manner as that of Example 3 with the exception that a mixture solution (1:5) of 49% hydrofluoric acid and hydrogen peroxid was used as an etchant. In two point six minutes after initialization of etching, the porous Si was sel ctively removed with th polycrystalline SI acting as an etch stopper, only the polycrystalline SI and the monocrystal fine SI being left behind, as shown in Fig. 3C.

## Example 14

Etching was performed in the same manner as that of Example 4 with the exception that a mixture solution (1:5) of 49% hydrofluoric acid and hydrogen peroxide was used as an etchant. In one point three minutes after initialization of etching, only the porous SI was selectively removed, only the polycrystalline SI and monocrystalline SI being left behind, as shown in Fig. 4B.

#### 15 Example 15

Etching was performed in the same manner as that of Example 5 with the exception that a mixture solution (1:5) of 49% hydrofluoric acid and hydrogen peroxide was used as an etchant. In this atching, only the porous Si was selectively removed, leaving the monocrystalline Si behind, as shown in Fig. 5C. Finally, the resist was removed (Fig. 5D).

## Example 16

Etching was performed in the same manner as that of Example 1 with the exception that a mixture solution (10:6:50) of 49% hydrofluoric acid, ethyl alcohol and hydrogen peroxide was used as an etchant. In twenty-six mixtures after initialization of etching, the porous Si was selectively removed with the remaining monocrystalline Si acting as an etch stooper, as shown in Fig. 19.

#### Example 17

Etching was performed in the same manner as that of Example 2 with the exception that a mixture solution (10:6:5) of 49% hydrofluoric acid, ethyl alcohol and hydrogen peroxide was used as an etchant. In one point four minutes after initialization of etching, the porous Si was selectively removed, only the monocrystalline SI being left behind, as shown in Fig. 28.

# Example 18

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Etching was performed in the same manner as that of Example 3 with the exception that a mixture solution (10:6:50) of 49% hydrofluoric acid, ethyl alcohol and hydrogen peroxide was used as an etchant. In two point eight minutes after initialization of etching, the porous Si was selectively removed with the polycrystalline SI acting as an etch stopper, only the polycrystalline SI and the monocrystalline SI being left behind, as shown in Fig. 3C.

# Example 19

Etching was performed in the same manner as that of Example 4 with the exception that a mixture solution (10:6:50) of 49% hydrofluoric acid, ettyl elcohol and hydrogen peroxide was used as an etchant. In one point four minutes after initialization of etching, the porous SI was selectively removed, only the polycrystalline SI and monocrystalline SI being left behind, as shown in Fig. 4B.

## Example 20

Etching was performed in the same manner as that of Example 5 with the exception that a mixture solution (100:6:50) of 49% hydrofluoric acid, ethyl alcohol and hydrogen peroxide was used as an etchant. In twenty-eight minutes after initialization of etching, the porous SI was selectively removed, only the monocrystalline SI being left behind, as shown in Fig. 5C. Finally, the resist was removed (Fig. 5D).

#### Example 21

Etching was performed in the same manner as that of Example 1 with th coeption that a buffered hydnoffur fic acid (NH<sub>4</sub>F: 36.2%, HF: 4.5%) was used as an tchant. In nin tennihutes after initialization of tching, the porous Si was a lectively removed with the monocrystalline Si acting as an etch stopp r, as shown in Fig. 18.

## Example 22

Etching was performed in the same manner as that of Example 2 with the exception that a buffered hydrofluoric acid (NHJF: 36.2%, HF: 4.5%) was used as an etchant. In seven seconds after initialization of etching, only the porous SI was selectively removed, leaving the monocrystalline SI behind, as shown in Fig. 2B.

#### Example 23

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Etching was performed in the same manner as that of Example 3 with the exception that a buffered hydrofluoric acid (NH,F: 3.6.2%, HF: 4.5%) was used as an etchant. In fourteen seconds after initialization of etching, the porous Si was selectively removed with the polycrystalline Si acting as an etch stopper, only the polycrystalline Si and the monocrystalline Si being left behind, as shown in Fig. 3C.

## Example 24

Etching was performed in the same manner as that of Example 4 with the exception that a buffered hydroftuoric acid (NH<sub>x</sub>F: 8.2%, HF: 4.5%) was used as an etchant. In seven seconds after initialization of etching, only the porous SI was selectively removed, leaving the polycrystalline SI and monocrystalline SI behind, as shown in Fig. 4B.

## Example 25

Etching was performed in the same manner as that of Example 5 with the exception that a buffered hydrofluoric acid (NH<sub>K</sub>F: 35.2%, HF: 4.5%) was used as an etchant. In nineteen minutes after initialization of etching, the porous SI was selectively removed, only the monocrystalline SI being left behind, as shown in FIg. 5C. Finally, the resist was removed (Fig. 5D).

## 55 Example 26

Etching was performed in the same manner as that of Example 1 with the exception that a mixture solution (10:1) of buffered hydrofluoric acid and ethyl alcohol was used as an etchant. In twenty-one minutes after initialization of etching, the porous SI was selectively removed with the monocrystalline SI acting as an etch stopper, only the monocrystalline SI being a left behind, as shown in Fig. 18.

## Example 27

Etching was performed in the same manner as that of Example 2 with the exception that a mixture solution (10:1) of buffered hydrofluoric acid and ethyl alcohol was used as an etchant. In seven seconds after initialization of etching, only the porous Si was selectively removed, leaving the monocrystalline Si behind, as shown in Fig. 2B.

# Example 28

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Etching was performed in the same manner as that of Example 3 with the exception that a mixture solution (10:1) of buffered hydrofluoric acid and ethyl alcohol was used as an etchant. In fourteen seconds after initialization of etching, the porous SI was selectively removed with the polycrystalline SI acting as an etch stopper, only the polycrystalline SI and the monocrystalline SI being left behind, as shown in Fig. 3C.

## Example 29

Etching was performed in the same manner as that of Example 4 with the xception that a mixture soluti n

(10:1) of buffered hydrofluoric acid and ethyl alcohol was used as an tchant. In a ven seconds after initialization of etching, only the porous Si was a lectively removid, leaving the polycrystallin. SI and minocrystalline Si behind, as shown in Fis. 48.

## 5 Example 30

Etching was performed in the same manner as that of Example 5 with the exception that a mixture solution (10:1) of buffered hydrofluoric acid and ethyl alcohol was used as an etchant. In twenty-one minutes after initialization of etching, the porous Si was selectively removed, only the monocrystalline SI being left behind, as shown in Fig. 5C. Finally, the resist was removed (Fig. 5D).

## Example 31

Etching was performed in the same manner as that of Example 1 with the exception that a mixture solution (1:5) of buffered hydrofluoric acid and hydrogen peroxide was used as an etchant. In nine minutes after initialization of etching, the porous SI was selectively removed with the monocrystalline SI acting as an etch stopper, only the monocrystalline SI being left behind, as shown in Fig. 1B.

#### Example 32

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Etching was performed in the same manner as that of Example 2 with the exception that a mixture solution (1:5) of buffered hydrofluoric acid and hydrogen peroxide was used as an etchant. In five seconds after initialization of etching, only the porous Si was selectively removed, leaving the monocrystalline Si behind, as shown in Fig. 2B.

## Example 33

Etching was performed in the same manner as that of Example 3 with the exception that a mixture solution (1:5) of buffered hydrofluoric acid and hydrogen peroxide was used as an etchant. In ten seconds after initialization of etching, the porous Si was selectively removed with the polycrystalline Si acting as an etch stopper, only the polycrystalline Si and the monocrystalline Si being left behind, as shown in Fig. 3C.

# Example 34

Etching was performed in the same manner as that of Example 4 with the exception that a mixture solution (1:5) of buffered hydrofluoric acid and hydrogen peroxide was used as an etchant. In five seconds after initialization of etching, only the porous Si was selectively removed, leaving the polycrystalline SI and monocrystalline SI behind, as shown in Fig. 48.

## Example 35

Etching was performed in the same manner as that of Example 5 with the exception that a mixture solution (1:5) of buffered hydrofluoric acid and hydrogen peroxide was used as an etchant. In nine minutes, the porous Si was selectively removed, only the monocrystalline Si being left behind, as shown in Fig. 5C. Finally, the resist was removed (Fig. 5D).

## Example 36

Etching was performed in the same manner as that of Example 1 with the exception that a mixture solution (106:50) of buffered hydrofluoric acid (NH<sub>4</sub>F: 36.2%, HF 4.5%), eithy alcohol and hydrogen peroxide was used as an etchant. In ten minutes after initialization of etching, the porous SI was selectively removed with the monocrystalline SI acting as an etch stopper, only the monocrystalline SI being left, as shown in Fig. 1B.

## Example 37

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Etching was performed in the same manner as that of Example 2 with the exception that a mixture solution (10:5:50) of buffered hydrofluorio acid (NH.F.: 36.2%, HF 4.5%), ethyl acohol and hydrogen peroxide was used as an etchant. In six seconds after initialization of tching, myth porous Si was selectively removed, leaving

the monocrystalline Si behind, as shown in Fig. 2B.

#### Example 38

Etching was perform d in the same manner as that of Exampi. 3 with the xception that a mixture is tution (10:6:50) of buffered hydrofluoric acid (NH<sub>4</sub>F: 36.2%, HF 4.5%), thyl alcohol and hydrogen peroxide was used as an etchant. In twelve seconds after initialization of etching, the porous SI was selectively removed with the polycrystalline SI acting as an etch stopper, only the polycrystalline SI and the monocrystalline SI being left behind, as shown in Fig. 3C.

## Example 39

Etching was performed in the same manner as that of Example 4 with the exception that a mixture solution (10:6:50) of buffered hydrofluoric acid (NH<sub>s</sub>F: 36.2%, HF 4.5%), ethyl alcohol and hydrogen peroxide was used as an etchant. In six seconds after initialization of etching, only the porous SI was selectively removed, leaving the polycrystalline SI and monocrystalline SI behind, as shown in Fig. 48.

#### Example 40

Etching was performed in the same manner as that of Example 5 with the exception that a mixture solution (10:6:50) of buffered hydrofluoric acid (NH<sub>2</sub>F: 38.2%, HF 4.5%), ethyl alcohol and hydrogen peroxide was used as netchant. In ten minutes after initialization of etching, the porous SI was selectively removed, only the monocrystalline SI being left behind, as shown in Fig. 5C. Finally, the resist was removed (Fig. 5D).

## 25 Example 41

Anodization was conducted on a P-type (100) monocrystalline SI substrate having a thickness of 200 μm in a 50% HF solution at a current density of 100 mAcrie?. The porous structure formation rate was 8.4 x and hence it took twenty four minutes for the 200 μm-thick P-yoe (100) Si substrate to be made entirely porous.

A Si epitaxial layer with a thickness of of 0.5 μm was grown on the P-type (100) porous Si substrate at a low temperature by molecular beam epitaxy (MBE). Deposition was conducted under the following conditions: Temperature: 700°C

Pressure: 1 x 10-9 Torr

Growth rate: 0.1 nm/sec.

Next, an optically polished fused silica glass substrate was placed on the surface of the epitaxial layer. The whole structure was then heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

Si<sub>3</sub>N<sub>4</sub>, was deposited to a thickness of 0.1 µm by plasma CVD method to cover the bonded substrates, and the nonly the nitried film on the porous substrate was removed by reactive ion etching. Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution while the solution was being stirred. In severnly eight minutes, the porous Si substrate was completely etched with the monocrystalline Si laver before as an etch stooper, only the monocrystalline Si laver before lief behind.

The etching rate of the non-porous monocrystalline Si was so low that only a maximum of 50 Å of non-porous monocrystalline Si was removed in seventy eight minutes. Since the ratio of the etching rate of the non-porous monocrystalline Si to that of the porous layer is 1:10° or more, the amount of non-porous layer which is etched (several tens angetroms) can be ignored in a practical operation. That is, the 200 µm-thick porous Si substrate was removed, and subsequently the Si<sub>2</sub>N<sub>4</sub> layer was removed with a result that the 0.5 µm-thick monocrystalline Si layer formed on the diass substrate remained.

The cross-section of the monocrystalline Si layer was observed with a transmission type electron microscope. It was found that no crystal defect was newly introduced in the Si layer and hence the Si layer had excellent crystalline structure.

#### Example 42

Anodization was conducted on a P type (100) monocrystalline SI substrate having a thickness of 200 µm in a 50% HF solution at a current density of 100 mA/cm². The porous structure formation rate was 8.4 µm/min and h noci it took twenty four minut sf r the 200 µm-thick Ptyp (100) SI substrate to be made nitiesty porous. A SI pitaxial layer with a thickness of 5.0 µm was grown on the Ptyp (100) porous SI substrate at a low tem-

p rature by plasma CVD. Depositi n was conducted und r th foll wing conditions:

Gas: SiH<sub>4</sub>
High-frequency power: 100 W
Temperatur: 800°C
Pressure: 1 x 10-2 Torr

Growth rat: 2.5 nm/sec.

Next, an optically polished glass substrate having a softening point of about 500 °C was placed on the surface of the epitaxial layer. The whole structure was heated at 450°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

 $Sl_3N_4$  was deposited to a thickness of 0.1  $\mu$ m by plasma CVD to cover the two bonded substrates, and then only the nitride film on the porous substrate was removed by reactive ion etching.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution while the solution was being stirred. In seventy eight minutes, the porous SI substrate was completely etched with the monocrystalline SI layer acting as an etch stopper, only the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in seventy eight minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous layer is 1: 10 for more, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed, and subsequently the Si<sub>2</sub>N<sub>4</sub> layer was removed with a result that the 5.0 µm-thick monocrystalline SI layer formed on the glass substrate having a low softening point remained.

# Example 43

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Anodization was conducted on a P type (100) monocrystalline Si substrate having a thickness of 200 µm in a 50% HF solution at a current density of 100 mA/cm². The porous structure formation rate was 8.4 µm/min and hence it took twenty four minutes for the 200 µm-thick P type (100) SI substrate to be made entirely porous. A SI epitaxial layer with a thickness of 1.0 µm was grown on the P type (100) porous SI substrate at a low temperature by blas sputtering. Deposition was conducted under the following conditions:

RF frequency: 100 MHz
High-frequency power: 600 W
Temperature: 300°C
Ar gas pressure: 8 x 10°3 Torr
Growth rate: 120 minutes
Target d. b. bias: -200 V

Target d.c. bias: 4200 V
Substrate d.c. bias: +5 V
Next, an optically polished glass substrate having a softening point of about 500 °C was placed on the surface of the epitaxial layer. The whole structure was heated at 450 °C in an oxygen atmosphere for 0.5 hours to

firmly join the two substrates to each other.

SI,N, was deposited to a thickness of 0.1 µm by plasma CVD to cover the two bonded substrates, and
then only the Initide film on the porous substrate was removed by reactive ion etching.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution while the solution was being stirred. In seventy eight minutes, the porous SI substrate was completely etched with the monocrystalline SI layer acting as an etch stopper, only the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in sevently leight minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous layer Is 1: 10 or more, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed, and subsequently the SI, N<sub>4</sub> layer was removed with a result that the 1.0 µm-thick monocrystalline SI layer on the glass substrate having a low softening point remained.

In case of coating of Aplexon Wax or Electron Wax in place of the SI<sub>3</sub>N<sub>4</sub> layer, the same effect was obtained and only the SI substrate made porous was completely removed.

#### Example 44

Anodization was conducted on a N type (100) monocrystalline Si substrate having a thickness of 200 µm in a 50% HF solution at a current density of 100 mA/cm². The porous structure formation rate was 8.4 µm/min and h noe it took twenty four minutes for the 200 µm-thick N type (100) SI substrat to be made nitr ly portion. A SI epitixal layer with a thickn so of 10 µm was grown on th N type (100) porous SI substrate at a low tem-

perature by liquid phas growth m thod under the following conditions:

Solvent: Sn, Solute:

Growth temperature: 900°C Growth atmosphere: H<sub>2</sub>

Growth time: 20 minut s

Next, an optically polished glass substrate having a soft ning point of about 800 °C was placed in the surface of the epitaxial layer. The whole structure was heated at 750°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

 $\mathrm{Si}_{n}\mathrm{N}_{4}$  was deposited to a thickness of 0.1  $\mu\mathrm{m}$  by plasma CVD to cover the two bonded substrates, and then only the nitride film on the porous substrate was removed by reactive ion etching.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution while the solution was being stirred. In seventy eight minutes, the prorus SI substrate was completely etched with the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in seventy eight minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous layer is 1: 10 or more, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed, and subsequently the SigN<sub>4</sub> layer was removed with a result that the 10 µm-thick monocrystalline SI layer on the glass substrate remained.

Coating of Apiezon Wax in place of the Si<sub>3</sub>N<sub>4</sub> layer was also effective and assured complete removal of only the porous SI substrate.

## Example 45

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Anodization was conducted on a P type (100) monocrystalline Si substrate having a thickness of 200 µm in a 50% HF solution at a current density of 100 mA/cm².

The porous structure formation rate was 8.4 µm/min and hence it took twenty four minutes for the 200 µmthick P type (100) S is ubstrate to be made entirely porous. A Si epitaxial layer with a thickness of 1.0 µmgrown on the P type (100) porous Si substrate at a low temperature by low-pressure CVD. Deposition was con-

ducted under the following conditions: Source gas: SiH<sub>4</sub> 800 SCCM Carrier gas: H<sub>2</sub> 150 liter/min

Temperature: 850°C Pressure: 1 x 10<sup>-2</sup> Torr

Growth rate: 3.3 nm/sec
Next, an optically polished fuzed silica glass substrate was placed on the surface of the epitaxial layer.
The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

Si<sub>3</sub>N<sub>4</sub> was deposited to a thickness of 0.1 µm by plasma CVD to cover the two bonded substrates, and then only the nitride film on the porous substrate was removed by reactive ion etching.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution will be the solution was being stirred. In severity eight minutes, the porous SI substrate was completely etched with the monocrystalline SI layer acting as an etch stopper, the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline Si was so low that only a maximum of 50 Å of non-porous monocrystalline Si was removed in seventy eight minutes. Since the ratio of the etching rate of the non-porous monocrystalline Si to that of the porous layer is 1: 10 or more, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous Si substrate was removed, and subsequently the Si,N<sub>4</sub> layer was removed with a result that the 1.0-thick µm monocrystalline Si layer on the silica glass substrate remained.

When SIH<sub>2</sub>Cl<sub>2</sub> was used as the source gas, the growth temperature had to be higher by several tens of degrees. However, high-speed etching characteristics to the porous substrate did not deteriorate.

#### Example 46

A Si epitaxial layer with a thickness of 1.0 μm was grown on a P type (100) Si substrate having a thickness of 200 μm by CVD. Deposition was conducted under the following conditions:

Reactiv gas flow rate: SiH<sub>4</sub> 1000 SCCM H<sub>2</sub> 230 liter/min

Temperature: 1080°C
Pressure: 760 Tom
Time: 2 min

Anodization was conducted on the substrat in a 50% HF solution at a current density of 100 mA/cm². The porous structure formation rate was 8.4 µm/min and hence it took tw nty four minutes for th 200 µm-thick Pype (100) 5 substrat to b made entirely porous. At that tim, there was no chang in the 5I epitaxial layer. Next, an optically polished fuzed silica glass substrate was placed on the surface of the epitaxial layer The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

Sl<sub>3</sub>N<sub>4</sub> was deposited to a thickness of 0.1 µm by plasma CVD to cover the two bonded substrates, and then only the nitride film on the porous substrate was removed by reactive ion etching.

Thereafter, selective stching was conducted on the bonded substrates in a 49% hydrofluorit cald solution while the solution was being stirred. In severity eight minutes, the porous SI substrate was completely removed with the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in seventy eight minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous leyer is 1: 10% the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed, and subsequently the SI<sub>N</sub>, layer was removed with a result that the 1.0 µm monocrystalline SI layer on the silica glass substrate remained.

The cross-section of the monocrystalline SI layer was observed with a transmission type electron microscope. It was found that no crystal defect was newly introduced in the SI layer and hence the SI layer had excellent crystalline structure.

## Example 47

A Si epitaxial layer with a thickness of 0.5 µm was grown on a P type (100) Si substrate having a thickness of 200 µm by CVD. Deposition was conducted under the following conditions:

Reactive gas flow rate: SiH<sub>2</sub>Cl<sub>2</sub> 1000 SCCM H<sub>2</sub> 230 liter/min

Temperature: 1080°C

Pressure: 80 Torr

Anodization was conducted on the substrate in a 50% HF solution at a current density of 100 mA/cm². The porous structure formation rate was 8.4 µm/min and hence it took twenty four minutes for the 200 µm-thick P type (100) SI substrate to be made entirely porous. As mentioned above, anodization made only the P type (100) SI substrate porous, and there was no change in the SI epitaxial layer.

Next, an optically pollshed fuzed silica glass substrate was placed on the surface of the epitaxial layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

 $\mathrm{Si}_{3}N_{4}$  was deposited to a thickness of 0.1  $\mu m$  by plasma CVD to cover the two bonded substrates, and then only the nitride film on the porous substrate was removed by reactive ion etching.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution was being stirred. In seventy eight minutes, the porous SI substrate was completely etched with the monocrystalline SI layer acting as an etch stopper, only the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in seventy eight minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous layer is 1: 10% the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed, and subsequently the SI<sub>0</sub>N<sub>4</sub> layer was removed with a result that the 0.5 µm-thick monocrystalline SI layer on the silica glass substrate remained.

The cross-section of the monocrystalline Si layer was observed by a transmission type electron microscope, it was found that no crystal defect was newly introduced in the Si layer and hence the Si layer had excellent crystalline structure.

## Example 48

A N typ Si lay r with a thickness of 1 µm was formed on a P type (100) SI substrate having a thickness

of 200 µm by proton implantation. Implantation rate of H+ was 5 x 1015 (ions/cm2).

Anodization was conducted on the substrate in a 50% HF solution at a current density of 100 mA/cm². The porous structure formation rate was 8.4 µm/min and hence it took twenty four minutes for the 200 µm-thick P type (100) Si substrate t b made entirely porous. As mentioned above, anodizatin made only the P type (100) Si substrate corous, and there was no chance in thin N typ. Si lay r.

Next, an optically polished fuzed silica glass substrat was placed on the surface of the N type SI layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

 $Sl_3N_4$  was deposited to a thickness of 0.1  $\mu m$  by plasma CVD to cover the two bonded substrates, and then only the nitride film on the porous substrate was removed by reactive ion etching.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution while the solution was being stirred. In seventy eight minutes, the porous SI substrate was completely etched with the monocrystalline SI laver acting as an etch stooper, only the monocrystalline SI laver being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in seventy eight minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous layer is 1:10°, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed, and subsequently the SI<sub>3</sub>N<sub>4</sub> layer was removed with a result that the 1 µm-thick monocrystalline SI layer on the silica glass substrate remained.

The cross-section of the monocrystalline Si layer was observed by a transmission type electron microscope. It was found that no crystal defect was newly introduced in the Si layer and hence the Si layer had excellent crystalline structure.

## Example 49

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Anodization was conducted on a P type (100) monocrystalline Si substrate having a thickness of 200 μm in a HF solution under the following conditions:

Applied voltage: 2.6 (V)

Current density: 30 (mA-cm<sup>-2</sup>)

Anodizing solution: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

Time: 1.6 (hour)

Thickness of porous Si: 200 (µm)

Porosity: 56 (%)

A Si epitaxial layer with a thickness of 0.5 μm was grown on the P type (100) porous Si substrate at a low temperature by molecular beam epitaxy (MBE). Deposition was conducted under the following conditions:

Temperature: 700°C

Pressure: 1 x 10<sup>-9</sup> Torr Growth rate: 0.1 nm/sec.

Next, a second SI substrate with a 5000 Å thick oxidized layer formed on the surface thereof was placed on the surface of the epitaxial layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution with the solution was being stirred, in sixty two minutes, the porous SI substrate was completely etched with the monocrystalline SI layer acting as an etch stopper, only the monocrystalline SI layer being left belind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in sixty-two minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous layer is 1: 10°, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed with a result that the 0.5 µm-thick monocrystalline SI layer formed on the SIO, layer remained.

The cross-section of the monocrystalline Si layer was observed by a transmission type electron microscope. It was found that no crystal defect was newly introduced in the Si layer and hence the Si layer had excellent crystalline structure.

## Example 50

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Anodization was conducted on a P type (100) monocrystalline SI substrate having a thickness of 200  $\mu m$  in a HF solution in the same manner as that of Example 49.

A Si epitaxial layer with a thickness of 0.5 µm was gr wn nth P typ (100) porous Si substrate at a low

temperature by plasma CVD. Deposition was c inducted under the following conditions:

Gas: SiH<sub>4</sub>
High-frequ ncy power: 100 W
T mperatur: 800°C
Pr ssur: 1 x 10-2 Torr

Growth rate: 2.5 nm/s c.

Next, a second Si substrate with a 5000 Å thick oxidized layer formed on the surface thereof was placed on the surface of the epitaxial layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution with the solution was being stirred. In skty two minutes, the portous SI substrate was completely etched with the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in sixty-two minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous layer is 1:10°, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed with a result that the 0.5 µm thick monocrystalline SI layer on the SIQ-gayer remained.

# Example 51

Anodization was conducted on a P type (100) monocrystalline SI substrate having a thickness of 200 μm in a HF solution in the same manner as that of Example 49.

A SI epitaxial layer with a thickness of 0.5 µm was grown on the P type (100) porous SI substrate at a low temperature by bias sputtering. Deposition was conducted under the following conditions:

RF frequency: 100 MHz
High-frequency power: 600 W
Temperature: 300°C
Ar gas pressure: 60 minutes
Growth time: 60 minutes
Target d.c. bias: -200 V
Substrate d.c. bias: +5 V

Next, a second Si substrate with a 5000 Å thick oxidized layer formed on the surface thereof was placed on the surface of the epitaxial layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acld solution will the solution was being stirred. In sixty two minutes, the porous SI substrate was completely etched with the monocrystalline SI layer acting as an etch stopper, the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in sixty-two minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous layer is 1:10°, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed with a result that the 0.5 µm thick monocrystalline SI layer on the SIO<sub>2</sub> layer remained.

#### Example 52

Anodization was conducted on a N type (100) monocrystalline Si substrate having a thickness of 200  $\mu m$  in a HF solution in the same manner as that of Example 49.

A SI epitaxial layer with a thickness of 5 µm was grown on the N type (100) porous SI substrate at a low temperature by liquid phase growth under the following conditions:

Solvent: Sn, Solute: 900°C
Growth atmosphere: H<sub>2</sub>
Growth rate: 10 minutes

Next, a second SI substrate with a 5000 Å thick oxidized layer formed on the surface thereof was placed on the surface of the epitaxial layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

SI

Th reaft r, s lectiv tching was conduct d n the bonded substrat s in a 49% hydrofluoric acid solution whil the solution was being stirred. In sixty two minutes, the porous Si substrate was compiletely etched with

the monocrystalline Si layer acting as an etch stopper, the monocrystalline Si layer being left behind.

The etching rate of the non-porous monocrystallin. Si was so low that Inly a maximum of 50 Å of non-porous monocrystalline Si was rem ved in sixty-two minutes. Sinc the ratio I th etching rate of the n n-porous monocrystallin. Si to that of th porous lay ris 1: 10°, th amount of non-porous layer which is etch if (several tens angetroms) can be ignored in a practical operation. That is, th 200 µm-thick porous SI substrat was removed with a result that th 5 µm thick monocrystalline SI layer or the SIOs (layer remained.

## Example 53

Anodization was conducted on a P type (100) monocrystalline SI substrate having a thickness of 200 μm in a HF solution in the same manner as that of Example 49.

A SI epitaxial layer with a thickness of 1.0 µm was grown on the P type (100) porous SI substrate at a low temperature by low-pressure CVD. Deposition was conducted under the following conditions:

Source gas: SiH<sub>4</sub>
Carrier gas: H<sub>2</sub>
Temperature: 850°C
Pressure: 1 x 10<sup>-2</sup> Torr

3.3 nm/sec

Next, a second Si substrate with a 5000 Å thick oxidized layer formed on the surface thereof was placed on the surface of the epitaxial layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly loin the two substrates to each other.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution with the solution was being stirred. In sixty two minutes, the porous SI substrate was completely etched with the monocrystalline SI layer scing as an etch stopper, the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in sixty-two minutes. Since the ratio of the etching rate of the non-porous monocrystalline SI to that of the porous layer is 1: 10°, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed with a result that the 1.0 µm thick monocrystalline SI layer on the SIO-) layer remained.

When SIH<sub>2</sub>Cl<sub>2</sub> was used as the source gas, the growth temperature had to be higher by several tens of degrees. However, high-speed etching characteristics to the porous substrate did not deteriorate.

# Example 54

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A Si epitaxial layer with a thickness of 1 μm was grown on a P type (100) Si substrate having a thickness of 200 μm by low-pressure CVD. Deposition was conducted under the following conditions:

Reactive gas flow rate: SiH<sub>2</sub>Cl<sub>2</sub> 1000 SCCM H<sub>2</sub> 230 liter/min

Temperature: 1080°C
Pressure: 80 Torr
Time: 2 min

Anodization was conducted on the substrate in a 50% HF solution at a current density of 100 mA/cm². The porous structure formation rate was 8.4 µm/min and hence it took twenty four minutes for the 200 µm-thick P type (100) SI substrate to be made entirely porous. As mentioned above, anodization made only the P type (100) SI substrate porous, and did not affect the SI epitaxial laver at all.

Next, a second SI substrate with a 5000 Å-thick oxidized layer formed on the surface thereof was placed on the surface of the epitaxial layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly loin the two substrates to each other.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution while the solution was being stirred. In skty two minutes, the porcus SI substrate was completely etched with the monocrystalline SI laver cating as an etch stopper, while the monocrystalline SI laver remained.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in sixty two minutes. Since the ratio of the etching rate of the non-porous noncrystalline SI to that of the porous layer is 1: 10°, the amount of non-porous tayer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed with a result that the 1.0 µm thick monocrystalline SI layer on the SIO, layer remained.

The cross-section of the monocrystalline Si layer was observed by a transmission type electron microscope. It was found that no crystal defect was newly introduced in the Si layer and hence the Si layer had excellent crystallin structure.

## Example 55

A SI epitaxial lay r with a thickn ss of 5 µm was grown on a P type (100) SI substrate having a thickn ss of 200 µm by atmosph ric CVD. Deposition was conducted under the following conditions:

Reactive gas flow rate: SiH<sub>2</sub>Cl<sub>2</sub> 1000 SCCM

H<sub>2</sub> 230 liter/min
Temperature: 1080°C

Pressure: 760 Torr

Anodization was conducted on the substrate in a HF solution in the same manner as that of Example 49. As mentioned above, anodization made only the P type (100) SI substrate porous, and did not affect the SI

Next, a second Si substrate with a 5000 Å-thick oxidized layer formed on the surface thereof was placed on the surface of the epitaxial layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0.5 hours to firmly join the two substrates to each other.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution with the solution was being stirred. In sixty two minutes, the prorus SI substrate was completely etched with the monocrystalline SI layer acting as an etch stopper, the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline Si was so low that only a maximum of 50 Å of non-porous monocrystalline Si was removed in sixty two minutes. Since the ratio of the etching rate of the non-porous monocrystalline Si to that of the porous layer is 1: 10°, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed with a result that the 5.0 µm thick monocrystalline Si layer on the SiQ layer remained.

The cross-section of the monocrystalline Si layer was observed by a transmission type electron microscope. It was found that no crystal defect was newly introduced in the Si layer and hence the Si layer had excellent crystalline structure.

# 30 Example 56

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A N type Si layer with a thickness of 1  $\mu$ m was formed on a P type (100) Si substrate having a thickness of 200  $\mu$ m by proton implantation. Implantation rate of H\* was 5 x 10<sup>15</sup> (lons/cm²).

Anodization was conducted on the substrate in a 50% HF solution at a current density of 100 mA/cm². The porous structure formation rate was 8.4 µm/min and hence it took twenty four minutes for the 200 µm-thick P type (100) SI substrate to be made entirely porous. As mentioned above, anodization made only the P type (100) SI substrate porous, and did not change the N type SI layer.

Next, a second SI substrate with a 5000 Å-thick oxidized layer formed on the surface thereof was placed on the surface of the N type SI layer. The whole structure was heated at 800°C in an oxygen atmosphere for 0,5 hours to firmly join the two substrates to each other.

Thereafter, selective etching was conducted on the bonded substrates in a 49% hydrofluoric acid solution will ethe solution was being stirred. In sixty two minutes, the porous SI substrate was completely etched with the monocrystalline SI layer acting as an etch stopper, the monocrystalline SI layer being left behind.

The etching rate of the non-porous monocrystalline SI was so low that only a maximum of 50 Å of non-porous monocrystalline SI was removed in sixty two minutes. Since the rate of the etching rate of the non-porous monocrystalline SI to that of the porous layer is 1: 10°, the amount of non-porous layer which is etched (several tens angstroms) can be ignored in a practical operation. That is, the 200 µm-thick porous SI substrate was removed with a result that the 1.0 µm thick monocrystalline SI layer on the SIO<sub>2</sub> layer remained.

The cross-section of the monocrystalline SI layer was observed by a transmission type electron microscope. It was found that no crystal defect was newly introduced in the SI layer and hence the SI layer had excellent crystalline structure.

## Example 57

A P-type (100) single-crystals (monocrystal) SI substrate of a thickness of 200  $\mu$  was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porcus structure formation rate then was about 8.4  $\mu$ m/min., and the P-type (100) SI substrate of a thickness of 200  $\mu$  was rendered porcus in its entirety for 24 minut s.

According to MBE (molecular beam epitaxy) method, an SI epitaxial lay r of 0.5 μ was grown at a lower temperature on th P-typ (100) porous SI substrate. The conditions for deposition are as follows;

temperature: 700 °C pressure: 1 x 10<sup>-9</sup> Torr

growth rate: 0.1 nm/sec.

Subsequ ntly, the surface of the pitaxial lay rewas thermally addized in a depth of 50 nm. A substrate of fused silica glass processed with optical polishing was bonded onto the thermally oxidized membran (i..., film), and both of the substrates were strongly bonded together by heating at 800 °C for 0.5 hour in oxygen atmosphere.

According to low pressure CVD, Sl<sub>3</sub>N<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane (film) on the porous substrate was removed by reactive ion etching.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid. In 78 minutes, only the single-crystal SI layer remainde without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal (monocrystal) with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the SI<sub>0</sub>N<sub>4</sub> layer, a single-crystal SI layer of a thickness of 0.5 µm was formed on the substrate of the silica glass.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

## Example 58

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A P-type (100) single-crystal Si substrate of a thickness of 200 µ was anodized in 50 % HF soulion. The current density then was 100 mA/cm². The porous structure formation rate then was abut 8.4 µm/min., and the P-type (100) Si substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes. According to plasma CVD method, an Si epitaxial layer of 5 µ was grown at a lower temperature on the P-type (100) porous Si substrate. The conditions for deposition are as follows;

gas: SiH<sub>4</sub>
high-frequency power: 100 W
temperature 800 °C

pressure: 1 x 10<sup>-2</sup> Torr growth rate: 2.5 nm/sec.

Subsequently, the surface of the epitaxial layer was thermally oxidized in a depth of 50 nm. A glass substrate, having being processed with optical polishing and having a softening point around 500 °C, was bonded onto the thermally oxidized membrane, and both of the substrates were strongly bonded together by heating at 450 °C for 0.5 hour in oxygen atmosphere.

According to plasma CVD method, Si<sub>3</sub>N<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid. In 78 minutes, only the single-crystal SI layer remained without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective rate of the ethcing rate of the porous layer to that of the non-porous SI single-crystal was as large as 10<sup>5</sup> or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the Si<sub>8</sub>N<sub>4</sub> layer, a single-crystal SI layer of a thickness of 5 µm was formed on the glass substrate of a lower softening point.

# Example 59

A P-type (100) single-crystal SI substrate of a thickness of 200 µ was anodized in 50 % HF solution. The current density then was 100 m/d/cm². The prorus structure formation rate then was abut 8.4 µm/min., and the P-type (100) SI substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes. According to thermal CVD method, an SI epitaxial layer of 5 µ was grown at a lower temperature on the P-type (100) porous SI substrate T n c ndfit in FT r dep sition are as follows:

gas: SiH<sub>4</sub>(0.6 Vmin), H<sub>2</sub> (100 Vmln)

temperature: 850 °C pressure: 50 Torr

growth rate; 0.1 µm/min.

Subs qu ntly, th surface of the pitaxial lay r was thermally oxidized in a d pth of 50 nm. A glass substrate, having b ing processed with optical polishing and having a s ftening point around 500 °C, was bonded onto the thermally oxidized membrane, and both of the substrates were strongly bonded tog ther by h ating at 450 °C for 0.5 hour in oxygen atmosphere.

at 450 °C for 0.5 nour in oxygen amouspiners.

According to plasma CVD method, SlyN<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid. In 78 minutes, only the single-crystal SI layer remained without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the Si<sub>2</sub>N<sub>4</sub> layer, a single-crystal SI layer of a thickness of 5 µm was formed on the glass substrate of a lower softening point.

A similar effect could be obtained by using Apiezon wax or electron wax, instead of Sl<sub>3</sub>N<sub>4</sub>, so that only the Si substrate rendered porous could be removed completely.

#### Example 60

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A P-type (100) single-crystal SI substrate of a thickness of 200 µ was anodized in 50 % HF solution. The terms thensity then was 100 mA/cm². The porous structure formation rate then was abut 8.4 µm/min., and the P-type (100) SI substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes. According to bias sputter method, an SI epitaxial layer of 1.0 µ was grown at a lower temperature on the P-type (100) porous SI substrate. The conditions for deposition are as follows;

RF frequency: 100 MHz

high-frequency power: 600 W temperature: 300 °C Ar gas pressure: 8 x 10-3 Torr growth time: 120 minutes target direct current bias: -200 V

substrate direct current blas: +5 V.

Subsequently, the surface of the epitaxial layer was thermally oxidized in a depth of 50 nm. A glass substrate, having being processed with optical polishing and having a softening point around 500 °C, was bonded onto the thermally oxidized membrane, and both of the substrate were strongly bonded together by heating at 450 °C for 0.5 hour in oxygen atmosphere.

According to plasma CVD method,  $Si_3N_4$  was deposited to 0.1  $\mu$ m, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid. In 78 minutes, only the single-crystal Si layer remained without etching, while the porous Si substrate was selectively etched with the single-crystal Si as a material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of thickness of 200 µ, rendered porous, was removed, and after the removal of the SI<sub>3</sub>N<sub>4</sub> layer, a single-crystal SI<sub>3</sub>Layer of a thickness of 1.0 µm was formed on the glass substrate of a lower melting point.

A similar effect could be obtained by using Aplezon wax or electron wax, instead of  $Sl_0N_4$ , so that only the SI substrate rendered porous could be removed completely.

## 5 Example 61

A N-type (100 single-crystal Si substrate of a thickness of 200  $\mu$  was anodized in 50 % HF solution. The current d nsity th n was 100 mA/cm<sup>2</sup>. Th porous structure formation rate th n was about 8.4  $\mu$ m/min., and

the N-typ (100) SI substrate of a thickn ss of 200 µ was rendered porous in its entirety for 24 minutes. According to liquid phas growth method, an SI pitaxial layer of 10 µ was grown at a lower timperature in the N-typ (100) porous Si substrat . The conditions for depositi n are as follows;

solvent Sn. Solute:

growth temp rature: 900°C

H<sub>2</sub> growth atmosphere:

growth time: 20 minutes.

Subsequently, the surface of the epitaxial layer was thermally oxidized in a depth of 50 nm. A glass substrate, having being processed with optical polishing and having a softening point around 800 °C, was bonded onto the thermally oxidized membrane, and both of the substrates were strongly bonded together by heating at 750 °C for 0.5 hour in oxygen atmosphere.

According to low pressure CVD method, Sl<sub>3</sub>N<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid. In 78 minutes, only the single-crystal SI layer remained without etching in 78 minutes, while the porous SI substrate was selectively etched with the single-crystal Si as a material for etching stopper and then completely removed.

The etching rate of the non-porous Si-single crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 105 or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the Si substrate, rendered porous, of a thickness of 200 µ was removed, and after the removal of the Si-N, layer, a single-crystal Si layer of a thickness of 10 µm was formed on the glass substrate of a lower softening point.

A similar effect could be obtained by using Apiezon wax or electron wax, instead of Si<sub>3</sub>N<sub>4</sub>, so that only the Si substrate rendered porous could be removed completely.

#### Example 62

According to CVD method, an Si epitaxial layer of 0.5 µ was grown at a lower temperature on a P-type (100) Si substrate of a thickness of 200 μ. The conditions for deposition are as follows:

reactive gas flow rate: SiH<sub>2</sub>Cl<sub>2</sub> 1000 SCCM

H 230 l/min.

temperature: 1080 °C pressure:

80 Torr

25

1 minute.

The present substrate was anodized in 50 % HF solution. The current density then was 100 mA/cm2. The porous structure formation rate then was about 8,4 μπ/min., and the P-type (100) Si substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes. As has been described above, the present anodization rendered only the P-type (100) Si substrate porous, but no change was observed in the Si epitaxial layer.

Subsequently, the surface of the epitaxial layer was thermally oxidized in a depth of 50 nm. A substrate of fused silica glass, processed with optical polishing, was bonded onto the thermally oxidized membrane, and both of the substrates were strongly bonded together by heating at 800 °C for 0.5 hour in oxygen atmosphere.

According to low pressure CVD method, Sl<sub>3</sub>N<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid. In 78 minutes, only the single-crystal Si layer remained without etching, while the porous Si substrate was selectively etched with the single-crystal Si as a material for etching stopper and then completely removed.

The etching rate of the non-porous Si single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 105 or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 μ, rendered porous, was removed, and after the removal of the SI<sub>3</sub>N<sub>4</sub> layer, a single-crystal SI layer of a thickness of 0.5 µm was formed on the glass substrate.

A similar effect could be obtained by using Aplezon wax or electron wax, instead of Si<sub>3</sub>N<sub>4</sub>, so that only the SI substrate rendered porous could be removed completely.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the SI layer and that excellent crystallinity was maintained.

## Example 63

A N-typ Si layer 1  $\mu$  was formed on the surface of a P-typ (100) Si substrat of a thickn ss of 200  $\mu$ , by

ion implantation of proton. The implanted amount of H\* was 5 x 10% lons/cm². Th substrate was anodized in 50 % HF solution. Th current density then was 100 mA/cm². Th porous structure formation rate th n was about 8.4 µm/mim., and the P-typ (100) Si substrate of a thickn as of 200 µ was rendered porous in its ntirely for 24 minutes. According to the pres nt anodization as has been described abov , nly th P-type (100) Si substrate was rendered porous, but no chang was bserved in the N-type Silayer. Subsequently, th surface of the N-typ single-crystal layer was thermally oxidized in a depth of 50 nm. A substrate of fused silica glass processed with optical polishing was bonded onto th thermally oxidized membran , and both of the substrates were strongly bonded tocether by heating at 880 °C for 0.5 hour in oxygen atmosphere.

According to low pressure CVD method, SI<sub>2</sub>N, was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive lon etching. Then, the bonded substrates were selectively etched with 49 % hydrofluoric ecid. In 78 minutes, only the single-crystal SI layer remained without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as Solution was extremely low, such as 50 angstroms or less even 78 inlustes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as  $10^6$  or more. The etched amount in non-porous leyer (several tens engstroms) is e practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of  $200\,\mu$ , rendered porous, was removed, and after the removal of the  $Si_2N_4$  layer, a single-crystal SI layer of a thickness of  $1.0\,\mu$ m was formed on the glass substrate.

A similar effect could be obtained by using Aplezon wax or electron wax, Insteed of Sl<sub>2</sub>N<sub>4</sub>, so that only the Si substrate rendered porous could be removed completely.

As a result of observation of the section under e transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

#### 25 Example 64

A P-type (100) single-crystal SI substrate of e thickness of 200  $\mu$  wes anodized in 50 % HF solution. The conditions for deposition are as follows;

applied voltage: 2.6 V

current density: 30 mA-cm<sup>-2</sup>

solution for anodization: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

time: 1.6 hours

thickness of porous Si: 200 µm

porosity: 56 %.

According to MBE method, en SI epitaxial leyer of 0.5  $\mu$  was grown et e lower temperature on the P-type (100) porous Si substrate. The conditions for deposition ere es follows;

temperature: 700 °C pressure: 1 x 10-9

growth rate: 0.1 nm/sec.

Subsequently, an oxidized layer of 1000 engstroms was formed on the surface of the epitaxial layer, and another Si substrate on the surface of which was formed an oxidized layer of 5000 angstroms, was bonded to the oxidized surface. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the Si substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid while under stirring. In 78 minutes, only the single-crystal Si layer remained without etching, while the porous Si substrate was selectively etched with the single-crystal Si as e material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as  $10^{9}$  or more. The etched emount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the SI<sub>N</sub>L, layer, a long-le-crystal SI layer of a thickness of 0.5 µm was formed on the SIO<sub>2</sub>. As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the SI layer and that excellent crystallinity was maintained.

## Example 65

A P-typ (100) single-crystal Si substrate of a thickness of 200 μ was anodized in HF solution.

The conditions for anodization ar as follows;

appli d voltage: current density: 2.6 V 30 mA.cm-2

HF:H2O:C2H5OH = 1:1:1 solution for anodization:

1.6 hours time:

thickn ss of porous Si:

10

15

200 um

porosity: 56 %

According to plasma CVD method, and SI epitaxial layer of 0.5 μ was grown at a lower temperature on the P-type (100) porous SI substrate. The conditions for deposition are as follows;

SiHz high-frequency power:

100 W 800 °C 1 x 10-2 Torr

temperature: pressure: growth rate: 2.5 nm/sec.

Subsequently, an oxidized layer of 1000 angstroms was formed on the surface of the epitaxial layer, and another SI substrate on the surface of which was formed as oxidized layer of 5000 angstroms was bonded to the oxidized surface. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the Si substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid while under stirring. In 78 minutes, only the single-crystal SI layer remained without etching, while the porous SI substrate was selectively etched with the single-crystal Si as a material for etching stopper and then completely removed.

The etching rate of the non-porous Si single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous Si single-crystal was as large as 105 or more. The etched amount in the non-porous layer (several tens anostroms) is a practically negligible decrease in membrane thickness. That is, the Si substrate of a thickness of 200 μ, rendered porous, was removed, and a single-crystal Si layer of a thickness of 0.5 um was formed on the SiO2.

#### Example 66

A P-type (100) single-crystal SI substrate of a thickness of 200 μ was anodized in HF solution.

The conditions for anodization are as follows;

applied voltage: 26 V

30 mA.cm<sup>-2</sup> current density:

thickness of porous SI:

HF:H2O:C2H5OH = 1:1:1 solution for anodization:

1 6 hours 200 um 56 %

porosity: According to bias sputter method, an SI epitaxial layer of 0.5  $\mu$  was grown at a lower temperature on the

P-type (100) porous SI substrate. The conditions for deposition are as follows; enn w

100 MHz RF frequency:

high-frequency power:

300 °C temperature: 8 x 10-3 Torr Ar gas pressure: 60 minutes growth time:

target direct current bias: -200 V

substrate direct current blas: +5 V

Subsequently, an oxidized layer of 1000 angstroms was formed on the surface of the epitaxial layer, and another Si substrate on the surface of which was formed an oxidized layer of 5000 angstroms was bonded to the oxidized surface. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the Si substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acis while under stirring. In 78 minutes, only the single-crystal SI layer remained without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and then completely removed. The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous Si single-crystal was as large as 105 or more. The etched amount in the non-porous layer (several tens angstroms) is a practically n gligible d crease in m mbrane thickness. That is, the Si substrate of a thick-

ness of 200  $\mu$ , r inder id porous, was removed, and a single-crystal SI layer of a thickness of 0.5  $\mu$ m was formed on the SiO2.

## Exampl 67

A N-typ (100) single-crystal SI substrat of a thickn ss of 200 μ was anodized in HF solution.

Th conditions for anodization ar as follows;

applied voltage: 2.6 V

current density: 30 mA.cm<sup>-2</sup> HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>2</sub>OH = 1:1:1

solution for anodization: 1.6 hours

200 µm thickness of porous Si:

porosity: 56 %

According to liquid phase growth method, an SI epitaxial layer of 5 µ was grown at a lower temperature on the N-type (100) porous Si substrate. The conditions for growth are as follows;

Sn, Solute: SI

900°C growth temperature: growth atmosphere: н,

growth period: 10 minutes

Subsequently, an oxidized layer of 1000 angstroms was formed on the surface of the epitaxial layer, and another Si substrate on the surface of which was formed an oxidized layer of 5000 angstroms was bonded to the oxidized surface. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the SI substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid while under stirring. Only the single-crystal Si layer remained without etching in 78 minutes, while the porous Si substrate was selectively etched with the single-crystal Si as a material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes layer, so that the selective ratio of the etching rate of the porous layer to that of the non-porous Si single-crystal was as large as 105 or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the Si substrate of a thickness of 200 µ, rendered porous, was removed, and a single-crystal SI layer of a thickness of 0.5 µm was formed on the SiO<sub>2</sub>.

#### Example 68

source gas:

A P-type (100) single-crystal Si substrate of a thickness of 200 μ was anodized in HF solution.

The conditions for anodization are as follows;

2.6 V applied voltage: current density: 30 mA-cm-2

HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1 solution for anodization:

time: 1.6 hours thickness of porous Si: 200 um

porosity: 56 %. SiH

According to low pressure CVD method, an Si epitaxial layer of 1.0 µ was grown at a lower temperature on the P-type (100) porous Si substrate. The conditions for deposition are as follows:

carrier gas: 850 °C temperature: 1 x 10-2 Torr pressure: 3.3 nm/sec. growth rate:

Subsequently, an oxidized layer of 1000 angstroms was formed on the surface of the epitaxial layer, and another SI substrate on the surface of which was formed an oxidized layer of 5000 angstroms was bonded to the oxidized surface. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the Si substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid while under stirring. In 78 minutes, only the single-crystal Si layer remained without etching in 78 minutes, while the porous SI substrate was selectively etched with the single-crystal Si as a material for etching stopper and then completely removed.

The etching rate of the non-porcus Si single-crystal with the etching solution was extremely low, such as

50 angstroms or I se av n 78 minut s later, so that the selectiv ratio of the tohing rat of th porous layer to that of th non-porous SI single-crystal was as larg as 10% or more. The toh d amount in th non-porous layer (several t ns angstroms) is a practically n glight decreas in m mbrane thickn ss. That is, th SI substrate, r nd red porous, of a thickness of 200 µ was r mov d, and a single-crystal SI layer of a thickness of 10 µm was formed on the SIQ. When SIN-CQ; was used a a source gas, it was required to rail of the provide temperature by sev rail tens of degrees. N v rithel ss, th accelerating tching characteristics to porous substrates was maintained.

#### Example 69

According to low pressure CVD method, an SI epitaxial layer of 1  $\mu$  was grown at a lower temperature on a P-type (100) porous SI substrate. The conditions for deposition are as follows;

reactive gas flow rate: SiH<sub>2</sub>Cl<sub>2</sub> 1000 SCCM H<sub>2</sub> 230 I/min

temperature: 1080 °C pressure: 80 Torr fime: 2 min.

The substrate was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous was 100 mA/cm². The porous curve formation rate then was 8.4 µm/min., and the P-type (100) SI substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes. According to the present anodization, as has been described above, only the P-type (100) SI substrate was rendered porous, but no change was observed in the SI epitaxial layer.

Subsequently, an oxidized layer of 1000 angstroms was formed on the surface of the epitaxial layer, and another Si substrate on the surface of which was formed an oxidized layer of 5000 angstroms was bonded to the oxidized surface. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the Si substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid while under stirring. In 78 minutes, only the single-crystal Si layer remained without etching, while the porous Si substrate was selectively etched with the single-crystal Si as a material for etching stopper and then completely removed.

The etching rate of the non-porous Si single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous Si single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and a single-crystal SI layer of a thickness of 1.0 µm was formed on the SiO<sub>2</sub>.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

## Example 70

temperature:

According to atmospheric pressure CVD method, an Si epitaxial layer of 5  $\mu$  was grown at a lower temperature on a P-type (100) Si substrate. The conditions for deposition are as follows;

reactive gas flow rate: SiH<sub>2</sub>Cl<sub>2</sub> 1000 SCCM

H<sub>2</sub> 230 l/min 1080 °C

pressure: 760 Torr

time: 1 min.
The Si substrate was anodized in HF solution.

The conditions for anodization are as follows; applied voltage: 2.6 V

current density: 30 mA-cm<sup>-2</sup> solution for anodization: HF:H<sub>2</sub>O:C<sub>2</sub>H<sub>5</sub>OH = 1:1:1

time: 1.6 hours
thickness of porous Si: 200 um

thickness of porous Si: 200 µ porosity: 56 %.

According to the present anodization as has been described above, only the P-type (100) SI substrate was rendered porous, but no change was observed in the Si epitaxial layer.

Subsequently, an oxidized layer of 1000 angstroms was formed on the surface f the epitaxial layer, and

another SI substrate on the surface of which was formed an xidiz d layer of 5000 angstroms was bond d to the oxidiz d surface. By heating in xygen atmosph r at 800 °C for 0.5 hour, both f th Si substrates w re strongly bond d tog th r.

Thin, thi bonded substrates were selectively etched with 49 % hydrofluoric acid while undir stirring. In 78 minutes, nly th single-crystal Si layer remained without tching, whill the porous SI substrat was selectively etch d with the single-crystal SI as a material for tching stopper and th n compl tely removed.

The tching rate of the non-porous SI single-crystal with the tching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous Si single-crystal was as large as 105 or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate, rendered porous, of a thickness of 200 µ was removed, and a single-crystal SI layer of a thickness of 5 um was formed on the SiO<sub>2</sub>. As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the SI layer and that excellent crystallinity was maintained.

#### Example 71

A N-type Si layer of 1  $\mu$  was formed on the surface of a P-type (100) Si substrate of a thickness of 200  $\mu$ , by ion implantation of proton. The implanted amount of H+ was 5 x 1015 ions/cm2. The substrate was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous structure formation rate then was 8.4 μm/min., and the P-type (100) Si substrate of a thickness of 200 μ was rendered porous in its entirety for 24 minutes. According to the present anodization as has been described above, only the P-type (100) SI substrate was rendered porous, but no change was observed in the N-type Si layer.

Subsequently, an oxidized layer of 1000 angstroms was formed on the surface of the N-type SI layer, and another SI substrate on the surface of which was formed an oxidized layer of 5000 angstroms was bonded to the oxidized surface. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the SI substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with 49 % hydrofluoric acid with stirring. In 78 minutes, only the single-crystal Si layer remained without etching, while the porous Si substrate was selectively etched with the single-crystal Si as a material for etching stopper and then completely removed.

The etching rate of the non-porous Si single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 78 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous Si single-crystal was as large as 105 or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the Si substrate, rendered porous, of a thickness of 200 µ was removed, and a single-crystal Si layer of a thickness of 1.0 µm was formed on SiO<sub>2</sub>.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

## Example 72

A P-type (100) single-crystal Si substrate of a thickness of 200 μ was anodized in 50 % HF solution. The current density then was 100 mA/cm2. The porous structure formation rate then was about 8.4 µm/min., and the P-type (100) Si substrate of a thickness of 200 μ was rendered porous in its entirety for 24 minutes.

According to MBE (monocular beam epitaxy) method, an Si epitaxlal layer was grown at a lower temperature on the P-type (100) porous Si substrate. The conditions for deposition are as follows;

700 °C temperature: pressure:

1 x 10-9 Torr

growth rate: 0.1 nm/sec.

Subsequently, a substrate of fused silica glass processed with optical polishing was bonded onto the the surface of the epitaxial layer, and both of the substrates were strongly bonded together by heating at 800 °C for 0.5 hour in oxygen atmosphere.

According to plasma CVD method, Si<sub>3</sub>N<sub>4</sub> was deposited to 0.1 μm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching. Then, the bonded substrates were selectively etching with a mixed solution of 49 % hydrofluoric acid and alcohol (10:1), without stirring. In 82 minutes, only the single-crystal Si layer remained without etching, while the porous Si substrate was selectively etched with the single-crystal Si as a material for etching stopper, and completely remov d.

The etching rate of th non-porous SI single-crystal with th etching solution was extremely low, such as 50 angstorms or less ven 82 minutes lat r, so that the selectiv ratio of th etching rate of th porou layer to that of th non-porous SI single-crystal was as large as 10° or more. The etch diamount in the non-porous layer (sev rail tens angstroms) is a practically negligible decreas in m mbrane thickn ss. That is, th SI substrate of a thickness of 200 µ, rendered porous, was r mov d, and after th removal of the SI<sub>5</sub>N<sub>4</sub> lay r, a single-crystal SI lay r of a thickn ss of 0.5 µm was formed on the substrat of the glass.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the SI layer and that excellent crystallinity was maintained.

#### Examples 73 to 86

The same procedure as in Examples 42 to 55 was effected, replacing the etching solution used in Examples 42 to 55 with that in Example 72. In any of the present Examples, consequently, a single-crystal SI layer was formed with extremely less crystal defect on insulating materials.

#### Example 87

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A N-type SI layer of 1  $\mu$  was formed on the surface of a P-type (100) Si substrate of a thickness of 200  $\mu$ , by ion implantation of proton. The implanted amount of H $^{*}$  was 5 x 10 $^{15}$  ions/cm<sup>2</sup>.

The substrate was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous structure formation rate then was 8.4µm/min., and the P-type (100) SI substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes. According to the present anodization as has been described above, only the P-type (100) SI substrate was rendered porous, but no change was observed in the N-type SI laver.

Subsequently, a second Si substrate on the surface of which was formed an oxidized layer of 5000 angstroms, was bonded to the surface of the N-type Si layer. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the Si substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with a mixed solution of 49 % hydrofluoric acid and alcohol (10:1) without stirring. In 82 minutes, only the single-crystal Si layer remained without etching, while the portous Si substrate was selectively etched with the single-crystal SI as a material for etching stopper and then completely removed.

The etching rate of the non-porous Si single-crystal with the etching solution was extremely low, such as 30 angstroms or less seven 82 minutes later, so that the selective ratio of the etching rate of the prorus layer to that of the non-porous Si single-crystal was as large as 10<sup>6</sup> or more. The etched amount in the non-porous layer (several tens angstrome) is a practically negligible decrease in membrane thickness. That is, the Si substrate of a thickness of 200 µ, rendered porous, was removed, and a single-crystal Si layer of a thickness of 1.0 µ m was formed on the SiOs layer.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

## Examples 88 to 102

The same procedure as in Examples 57 to 71 was effected, replacing the etching solution in Examples 57 to 71 with that in Examples 72. In any of the present Examples, consequently, a single-crystal Si layer was formed with extremely less crystal defect on insulating insterials.

#### Example 103

A P-type (100) single-crystal SI substrate of a thickness of 200 μ was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous structure formation rate then was about 8.4 μm/min., and the P-type (100) SI substrate of a thickness of 200 μ was rendered porous in its entirely for 24 minutes.

According to MBE (molecular beam epitaxy) method, an SI epitaxial layer of  $0.5 \,\mu$  was grown at a lower temperature on the P-type (100) porous SI substrate. The conditions for deposition are as follows; temperature:  $700 \,^{\circ}\text{C}$ 

pressure: 1 x 10<sup>-9</sup> Torr

growth rate: 0.1 nm/sec.

Subsequently, a substrate of fused silica glass processed with optical polishing was bonded onto the surface of the pitaxial layer, and both of the substrates were strongly bond d together by heating at 800 °C for

0.5 hour in oxygen atmosph re.

According to plasma CVD method, SlyN, was deposited at 0.1 µm, th r by coating th bonded two substrates. Th reaft r, nly th nitrid membran on th porous substrate was removed by reactiv ion tothing. Then, the bonded substrates were selectively tot d with a mixed solution of 49 % hydrofluoric acid and aqueous hydrog n peroxid solution (1:5), while under stiming. In 62 minutes, only the single-crystal SI layer remain d without kning, whill the porous SI substrate was a I ctiv ly etched with th single-crystal SI as a material for etching stopper and completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 62 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the SI<sub>5</sub>N<sub>6</sub> layer, a single-crystal SI layer of a thickness of 0.5 µm was formed on the substrate of the glass.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the SI layer and that excellent crystallinity was maintained.

## Examples 104 to 118

The same procedure as in Examples 42 to 56 was effected, replacing the etching solution in Examples 42 to 56 with that in Examples 103. In any of the present Examples, consequently, a single-crystal Si layer was formed with extremely less crystal defect on insulating materials.

#### Example 119

A P-type (100) single-crystal Si substrate of a thickness of 200 µ was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous structure formation rate then was about 8.4 µm/min., and the P-type (100) Si substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes.

According to MBE (molecular beam epitaxy) method, an SI epitaxial layer of  $0.5~\mu$  was grown at a lower temperature on the P-type (100) porous SI substrate. The conditions for deposition are as follows;

temperature: 700 °C

pressure: 1 x 10 ° Torr growth rate: 0.1 nm/sec.

Subsequently, the surface of the epitaxial layer was thermally oxidized in a depth of 50 nm. A substrate of fused silica glass processed with optical polishing was bonded onto the thermally oxidized membrane, and both of the substrates were strongly bonded together by heating at 800 °C for 0.5 hour in oxygen atmosphere.

According to low pressure CVD method, SI<sub>3</sub>N<sub>4</sub> was deposited to 0,1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching.

Then, the bonded substrates were selectively etched with a mixed solution of 49 % hydrofluoric acid and aqueous hydrogen peroxide solution (1:5). In 62 minutes, only the single-crystal SI layer remained without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and completely removed.

The etching rate of the non-porous Si single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 62 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous Is ingle-crystal was as large as 10<sup>6</sup> or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the Si substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the SigN<sub>k</sub> layer, a single-crystal Si layer of a thickness of 0.5 µm was formed on the substrate of the silica class.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

## Examples 120 to 133

The same procedure as in Examples 58 to 71 was effected, replacing the etching solution in Examples 58 to 71 with that in Example 103. In any of the present Examples, consequently, a single-crystal SI layer was formed with extremely less crystal defect on insulating materials.

## Example 134

A P-typ (100) single-crystal Si substrate of a thickness of 200 μ was anodized in 50 % HF solution. Th curr ntd nsity then was 100 mA/cm². Th porous structure formation rat th n was 3.4 μπ/mln, and th P-type (100) Si substrat of a thickness of 200 μ was rendered porous in its ntirety for 24 minutes.

According to MBE (mol cular b am pitaxy) method, an SI epitaxial layer of 0.5 μ was grown at a lower temperature on th P-type (100) porous SI substrate. The conditions for diposition are as follows;

temperature: 700 °C pressure: 1 x 10-9 Torr

growth rate: 0.1 nm/sec.

Subsequently, a substrate of fused silica glass processed with optical polishing was bonded onto the surface of the epitaxial layer, and both of the substrates were strongly bonded together by heating at 800 °C for 0.5 hour in oxygen atmosphere.

According to plasma CVD method, Sl<sub>2</sub>N<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching. Then, the bonded substrates were selectively etched with a mixed solution of 49 % hydroduroir acid, alcohol and aqueous hydrogen peroxide solution (10:6:50), without stirring. In 65 minutes, only the single-crystal SI layer remained without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and completely removed.

The etching rate of the non-porous Si single crystal with the etching solution was extremely low, such as approximately slightly less than 40 angstroms even 65 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous Si single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the Si substrate of a thickness of 200  $\mu$ , rendered porous, was removed, and after the remote of the SiN<sub>4</sub> layer, a single-crystal Si layer of a thickness of 0.5  $\mu$ m was formed on the substrate of the silical state.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

#### 30 Examples 135 to 148

The same procedure as in Examples 42 to 55 was effected, replacing the etching solution in Examples 42 to 55 with that in Example 134. In any of the present Examples, consequently, a single-crystal Si layer was formed with extremely less crystal defect on insulating materials.

## Example 149

A N-type SI layer of 1 μ was formed on the surface of a P-type (100) SI substrate of a thickness of 200 μ, by ion implantation of proton, The implanted amount of H<sup>+</sup> was 5 x 10<sup>15</sup> ions/cm<sup>2</sup>.

The substrate was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous structure formation rate then was 8.4 µm/mlm., and the P-type (100) SI substrate of a thickness c020 µ was rendered porous in its entirety for 24 minutes. According to the present anodization as has been described above, only the P-type (100) SI substrate was rendered porous, but no change was observed in the N-type SI laver.

Subsequently, a second SI substrate on the surface of which was formed an oxidized layer of 5000 angstroms was bonded to the surface of the N-type SI layer. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with a mixed solution of 49 % hydrofluoric acid, alcohol and aqueous 30 % hydrogen peroxide solution (10:6:50) without stirring. In 65 minutes, only the single-crystal SIIayer remained without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper, and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 65 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and a single-crystal SI layer of a thickness of 1.0 µm was formed on the Sicy layer.

As a result of observation of the section under a transmission-type of ctron microscope, it was confirmed

that no new crystal defect was introduced in the Si layer and that excell int crystallinity was maintained.

#### Examples 150 to 164

Th same procedure as in Exampl s 57 to 71 was effected, replacing th tching solution in Examples 57 to 71 with that in Exampl s 34. In any of th pr sent Exampl s, consequ ntly, a single-crystal SI lay r was formed with externely less crystal defect on insulating materials.

#### Example 165

A P-type (100) single-crystal SI substrate of a thickness of 200 µ was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous structure formation rate then was 8.4 µm/min., and the P-type (100) SI substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes.

According to MBE (molecular beam epitaxy) method, an SI epitaxial layer of  $0.5 \mu$  was grown at a lower temperature on the P-type (100) porous SI substrate. The conditions for deposition are as follows;

temperature: 700 °C
pressure: 1 x 10-9 Torr

growth rate; 0.1 nm/sec.

Subsequently, a substrate of fused silica glass processed with optical polishing was bonded onto the surface of the epitaxial layer, and both of the substrates were strongly bonded together by heating at 800 °C for 0.5 hour in oxygen atmosphere.

According to plasma CVD method, Sl<sub>3</sub>N<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching. Then, the bonded substrates were selectively etched, under stirring, with buffered hydrofluoric acid. In 258 minutes, only the single-crystal Slayer remained without etching, while the porous Slaubstrate was selectively etched with the single-crystal Slayer and there are substrated with the single-crystal Slayer and the removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 100 angstroms or less even 258 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10<sup>6</sup> or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the SigN<sub>4</sub> layer, a single-crystal SI layer of a thickness of 0.5 µm was formed on the substrate of the glass.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the SI layer and that excellent crystallinity was maintained.

## Examples 166 to 180

The same procedure as in Examples 42 to 56 was effected, replacing the etching solution in Examples 42 to 56 with that in Example 185. In any of the present Examples, consequently, a single-crystal SI layer was formed with extremely less crystal defect on insulating materials.

## Example 181

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A P-type (100) single-crystal SI substrate of a thickness of 200 μ was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous structure formation rate then was about 8.4 μm/min., and the P-type (100) SI substrate of a thickness of 200 μ was rendered porous in its entirety for 24 minutes.

According to MBE (molecular beam epitaxy) method, an Si epitaxial layer of  $0.5~\mu$  was grown at a lower temperature on the P-type (100) porous SI substrate. The conditions for deposition are as follows;

temperature: 700 °C pressure: 1 x 10-9 Torr

pressure: 1 x 10<sup>-8</sup> Torr growth rate: 0.1 nm/sec.

Subsequently, the surface of the epitaxial layer was thermally oxidized in a depth of 50 nm. A substrate of fused silica glass processed with optical polishing was bonded onto the thermally oxidized membrane, and both of

According to low pressure CVD method,  $Sl_3N_4$  was deposited to 0.1  $\mu$ m, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching.

Then, the bonded substrates were immersed in buffered hydrofluoric acid, and stirred. In 258 minutes, only the single-crystal Si layer remained without etching, while the porous Si substrates was selectively etched with

the single-crystal SI as a material for etching stopp r and then completely removed.

The tching rate of the non-porous Si ingle-crystal with the tching solution was extremely low, such as 100 angstroms or less ven 258 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the n n-porous Si single-crystal was as larg as 105 or more. The tiched amount in the non-porous layer (several t ns angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and aft r the removal of the Si<sub>3</sub>N<sub>4</sub> lay r, a sing-I -crystal Si layer of a thickn ss of 0.5 μm n th substrate of the silica glass was formed.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the SI layer and that excellent crystallinity was maintained.

## Examples 182 to 195

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The same procedure as in Examples 58 to 71 was effected, replacing the etching solution in Examples 58 to 71 with that in Example 165. In any of the present Examples, consequently, a single-crystal SI layer was formed with extremely less crystal defect on insulating materials.

## Example 198

A P-type (100) single-crystal SI substrate of a thickness of 200 μ was anodized in 50 % HF solution. The current density then was 100 mA/cm2. The porous structure formation rate then was 8.4 µm/min., and the P-type (100) Si substrate of a thickness of 200 μ was rendered porous in its entirety for 24 minutes.

According to MBE (molecular beam epitaxy) method, an Si epitaxial layer of 0.5 μ was grown at a lower temperature on the P-type (100) porous Si substrate. The conditions for deposition are as follows;

700 °C temperature: 1 x 10-9 Torr

pressure:

0.1 nm/sec. growth rate:

Subsequently, a substrate of fused silica glass processed with optical polishing was bonded onto the surface of the epitaxial layer, and both of the substrates were strongly bonded together by heating at 800 °C for 0.5 hour in oxygen atmosphere.

According to plasma CVD method, Si<sub>3</sub>N<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching. Then, the bonded substrates were selectively etched in a mixed solution of buffered hydrofluoric acid and alcohol (10:1) without stirring. In 275 minutes, only the single-crystal Si layer remained without etching, while the porous Si substrate was selectively etched with the single-crystal Si as a material for etching stopper and completely removed.

The etching rate of the non-porous Si single-crystal with the etching solution was extremely low, such as 100 angstroms or less even 275 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 105 or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the Si<sub>3</sub>N<sub>4</sub> layer, a singie-crystal Si layer of a thickness of 0.5 μm was formed on the substrate of the glass.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

#### Examples 197 to 210

The same procedure as in Examples 42 to 55 was effected, replacing the etching solution in Examples 42 to 55 with that in Example 196. In any of the present Examples, consequently, a single-crystal SI layer was formed with extremely less crystal defect on insulating materials.

## Example 211

A N-type SI layer of thickness of 1  $\mu$  was formed on the surface of a P-type SI layer of 200  $\mu$  in thickness, by Ion Implantation of proton, The implanted amount of H+ was 5 x 1015 lons/cm2.

The substrate was anodized in 50 % HF solution. The current density then was 100 mA/cm<sup>2</sup>. The porous structure formation rate then was 8.4  $\mu$ m/min., and the N-type (100) Si substrate of a thickness of 200  $\mu$  was rendered porous in its entirety for 24 minutes. According to the present anodization as has been described abov, only the P-typ (100) Si substrat was rendered porous, but no change was observed in the N-type Si iayer.

Subs qu ntly, a second SI substrat nth surface of which was formed an oxidized lay r of 5000 angstroms was bonded the surface of the N-type SI lay r. By heating in oxygen atmosph re at 800 °C for 0.5 hour, b th of the substrates were strongly bonded togeth r.

Then, the bind disubstrates were selectively etched with a mixed solution of buffered hydrofluoric acid (HF:4.46 %, NH,F:36.2 %) and ethyl alcohol (10:1), without stirring. In 275 minutes, only the single-crystal SI layer mained without tching, while the porous SI substrate was silectivity etchind with the single-crystal SI as a material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as approximately slightly less than 40 angstroms even 275 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and a single-crystal SI layer of a thickness of 1.0 µm was formed on the SIO<sub>2</sub> layer.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the SI layer and that excellent crystallinity was maintained.

#### Examples 212 to 226

The same procedure as in Examples 57 to 71 was effected, replacing the etching solution in Examples 57 to 71 with that in Examples 198. In any of the present Examples, consequently, a single-crystal Si layer was formed with extremely less crystal defect on insulating materials.

## Example 227

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A P-type (100) single-crystal SI substrate of a thickness of 200 μ was anodized in 50% HF solution. The current density then was 100 mA/cm². The prorus structure formation rate then was 8.4 μm/min., and the P-type (100) SI substrate of a thickness of 200 μ was rendered porous in its entirety for 24 minutes.

According to MBE (molecular beam epitaxy) method, an Si epitaxial layer of  $0.5~\mu$  was grown at a lower temperature on the P-type (100) porous Si substrate. The conditions for deposition are as follows;

temperature: 700 °C pressure: 1 x 10-9 Torr

growth rate: 0.1 nm/sec.

Subsequently, a substrate of fused silica glass processed with optical polishing was bonded onto the surface of the epitaxial layer, and both of the substrates were strongly bonded together by heating at 800 °C for 0,5 hour in oxygen atmosphere.

According to plasma CVD method, Si<sub>b</sub>N<sub>a</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive lon etching. Then, the bonded substrates were selectively etched in a mixed solution of buffered hydrofluoric acid and aqueous hydrogen peroxide solution (1:5) with stirring. In 190 minutes, only the single-crystal Si layer remained without etching, while the porous Si substrate was selectively etched with the single-crystal Si as a material for etching stopper, and completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 50 angstroms or less even 190 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous Isingle-crystal was as large as 10% or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the SI<sub>N</sub>4 layer, a single-crystal SI layer of a thickness of 0.5 µ m was formed on the substrate of the silica glass.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

## Examples 228 to 242

The same procedure as in Examples 42 to 56 was effected, replacing the etching solution in Examples 42 to 56 with that in Example 227. In any of the present Examples, consequently, a single-crystal SI layer was formed with extremely less crystal defect on insulating materials.

## Example 243

A P-type (100) single-crystal SI substrat of a thickness of 200 µ was anodized in 50% HF solution. The current of naity th n was 100 mA/cm?. Th porous structure formation rate th n was about 8.4 µm/min, and the P-type (100) SI substrate of a thickness of 200 µ was rend red porous in its nitrety for 24 minutes.

According to MBE (mol cular b am epitaxy) method, an SI pitaxlal layer of 0.5  $\mu$  was grown at a lower temperature on th P-typ (100) porous SI substrate. The conditions for d position ar as follows;

temperature: 700 °C

pressure: 1 x 10<sup>-8</sup> Tor growth rate: 0.1 nm/sec.

subsequently, the surface of the epitaxial layer was thermally oxidized in a depth of 50 mm. A substrate of fused silica glass processed with optical polishing was bonded onto the thermally oxidized membrane, and both of the substrates were strongly bonded together by heating at 800 °C for 0.5 hour in oxygen atmosphere.

According to low pressure CVD method, SI<sub>3</sub>N<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive ion etching.

Then, the bonded substrates were immersed in a mixed solution of buffered hydrofluoric acid and aqueous hydrogen peroxide solution (1:5), and stirred. In 190 minutes, only the single-crystal SI layer remained without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as 70 angstroms or less even 190 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrate of a thickness of 200 µ, rendered porous, was removed, and after the removal of the Si<sub>1</sub>N<sub>4</sub> layer, a single-crystal SI layer of a thickness of 0.5 µm was formed on the substrate of the silical glass.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the SI layer and that excellent crystallinity was maintained.

#### Examples 244 to 257

Examples 244 to 20

The same procedure as in Examples 58 to 71 was effected, replacing the etching solution in Examples 58 to 71 with that in Example 243. In any of the present Examples, consequently, a single-crystal SI layer was formed with extremely less crystal defect on insulating materials.

# 35 Example 258

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A P-type (100) single-crystal Si substrate of a thickness of 200 µ was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous structure formation rate then was 8.4 µm/min., and the P-type (100) Si substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes.

According to MBE (molecular beam epitaxy) method, an Si epitaxial layer of  $0.5~\mu$  was grown at a lower temperature on the P-type (100) porous Si substrate. The conditions for deposition are as follows;

temperature: 700 °C
pressure: 1 x 10-9 Torr
growth rate: 0.1 nm/sec.

Subsequently, a substrate of fused silica glass processed with optical polishing was bonded onto the surface of the epitaxial layer, and both of the substrates were strongly bonded together by heating at 800 °C for 0.5 hour in oxygen atmosphere.

According to plasma CVD method, SI<sub>3</sub>N<sub>4</sub> was deposited to 0.1 µm, thereby coating the bonded two substrates. Thereafter, only the nitride membrane on the porous substrate was removed by reactive lone tiching. Then, the bonded substrates were selectively etched in a mixed solution of buffered hydrofluoric acid, alcohol and aqueous hydrogen peroxide solution (10:6:50) without stirring, in 205 minutes, only the single-crystal SI layer remained without stirring, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and then completely removed.

The etching rate of the non-porous SI single-crystal with the etching solution was extremely low, such as asymmetry slightly less than 40 angstroms even 205 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10<sup>s</sup> or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the SI substrat of a thickness of 200 µ, rend red porous, was removed, and after the remove

of the Sl<sub>3</sub>N<sub>4</sub> layer, a single-crystal SI layer of a thickness of 0.5 μm was formed on the substrate of the silica

As a result of observation of the section undir a transmission-type lectron microscope, it was confirm of that no new crystal deficit was introduced in the SI layer and that ix xxell in transmission-type.

#### Exampl s 259 to 272

The same procedure as in Examples 42 to 55 was effected, replacing the etching solution in Examples 42 to 55 with that in Example 258. In any of the present Examples, consequently, a single-crystal SI layer was formed with extremely less crystal defect on insulating materials.

#### Example 273

A N-type SI layer of thickness of 1  $\mu$  was formed on the surface of a P-type (100) SI substrate of thickness of 200  $\mu$ , by ion implantation of proton. The implanted amount of H<sup>+</sup> was 5 x 10<sup>15</sup> lons/cm<sup>2</sup>.

The substrate was anodized in 50 % HF solution. The current density then was 100 mA/cm². The porous structure formation rate then was 8.4 µm/min., and the P-type (100) Si substrate of a thickness of 200 µ was rendered porous in its entirety for 24 minutes. According to the present anodization as has been described above, only the P-type (100) Si substrate was rendered porous, but no change was observed in the N-type SI

Subsequently, a second Si substrate on the surface of which was formed an oxidized layer of 5000 angstroms was bonded to the surface of the N-type Si layer. By heating in oxygen atmosphere at 800 °C for 0.5 hour, both of the substrates were strongly bonded together.

Then, the bonded substrates were selectively etched with a mixed solution of buffered hydrofluoric acid (H:F.4.6 %, NH<sub>c</sub>F:38.2%), ethyl alcohol and 30% aqueous hydrogen peroxide solution (10.6:50), without stiring. In 180 minutes, only the single-crystal Si layer remained without etching, while the porous SI substrate was selectively etched with the single-crystal SI as a material for etching stopper and then completely removed.

The etching rate of the non-porous Si single-crystal with the etching solution was extremely low, such as approximately slightly less than 40 angstroms even 180 minutes later, so that the selective ratio of the etching rate of the porous layer to that of the non-porous SI single-crystal was as large as 10° or more. The etched amount in the non-porous layer (several tens angstroms) is a practically negligible decrease in membrane thickness. That is, the Si substrate of a thickness of 200 µ, rendered porous, was removed, and a single-crystal SI layer of a thickness of 1.0 µm was formed on the SiO<sub>2</sub> layer.

As a result of observation of the section under a transmission-type electron microscope, it was confirmed that no new crystal defect was introduced in the Si layer and that excellent crystallinity was maintained.

#### Examples 274 to 288

The same procedure as in Examples 57 to 71 was effected, replacing the etching solution in Examples 57 to 71 with that in Example 27. In any of the present Examples, consequently, a single-crystal SI layer was formed with externely less crystal defect on insulating materials.

## Claims

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- 1. A chemical etching solution for porous Si comprising hydrofluoric acid as etchant.
- A chemical etching solution for porous Si comprising a mixture of hydrofluoric acid and an alcohol as etchant.
- A chemical etching solution for porous SI comprising a mixture of hydrofluoric acid and hydrogen peroxide
  as etchant.
  - A chemical etching solution for porous Si comprising a mixture of hydrofluoric acid, an alcohol and hydrogen peroxide as etchant.
  - 5. A chemical etching solution for porous Si comprising buffered hydrofluoric acid as etchant.

- A chemical etching solution for porous Si comprising a mixture of buffered hydrofluoric acid and an alcohol
  as tchant.
- A chemical etching solution for porous SI comprising a mixture of buffered hydrofluoric acid and hydrog n
  p roxide as tchant.
  - A chemical etching solution for porous SI comprising a mixture of buffered hydrofluoric acid, an alcohol
    and hydrogen peroxide as etchant.
- 9. A chemical etching method for porous Si, which comprises using hydrofluoric acid as etchant.
  - A chemical etching method for porous SI, which comprises using a mixture of hydrofluoric acid and an alcohol as etchant.
  - A chemical etching method for porous SI, which comprises using a mixture of hydrofluoric acid and hydrogen peroxide as etchant.
  - A chemical etching method for porous SI, which comprises using a mixture of hydrofluoric acid, an alcohol
    and hydrogen peroxide as etchant.
- 13. A chemical etching method for porous Si, which comprises using buffered hydrofluoric acid as etchant.
  - 14. A chemical etching method for porous Si, which comprises using a mixture of buffered hydrofluoric acid and an alcohol as etchant.
  - A chemical etching method for porous SI, which comprises using a mixture of buffered hydrofluoric acid and hydrogen peroxide as etchant.
    - A chemical etching method for porous Si, which comprises using a mixture of buffered hydrofluoric acid, an alcohol and hydrogen peroxide as etchant.
- 17. A method for preparing a semiconductor member, which comprises:

forming a substrate having a non-porous silicon monocrystalline layer and a porous silicon layer, bonding another substrate having a surface made of an insulating material to the surface of said monocrystalline layer, and

etching to remove said porous silicon layer by immersing in hydrofluoric acid.

- 18. A method for preparing a semiconductor member, which comprises:
  - forming a substrate having a non-porous silicon monocrystalline layer and a porous silicon layer, bonding another substrate having a surface made of an insulating material to the surface of said monocrystalline layer; and
- etching to remove said porous silicon layer by immersing in a mixture of hydrofluoric acid and an alcohol.
  - 19. A method for preparing a semiconductor member, which comprises:
- forming a substrate having a non-porous silicon monocrystalline layer and a porous silicon layer, bonding another substrate having a surface of an insulating material to the surface of said monocrystalline layer, and
  - etching to remove said porous silicon layer by immersing in a mixture of hydrofluoric acid and hydrogen peroxide.
- 20. A method for preparing a semiconductor member, which comprises:

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- forming a substrate having a non-porous silicon monocrystalline layer and a porous silicon layer, bonding another substrate having a surface made of an insulating material to the surface of said monocrystalline layer; and
- etching to remove said porous silicon layer by immersing in a mixture of hydrofluoric acid, an alcohol and hydrogen peroxide.
- 21. A method for preparing a semiconductor member, which comprises:

forming a substrate having a non-porous silicon monocrystalline layer and a porous silicon layer;

bonding another substrate having a surface mad of an insulating material to the surface of said monocrystallin layer, and

etching to remov said porous afficon lay r by immersing in buff red hydrofluoric acid.

22. A method for preparing a semiconductor member, which comprises:

forming a substrate having a non-porous sillcon monocrystallin lay r and a porous sillcon lay r, bonding another substrate having a surface made of an insulating material to the surface of said monocrystalline laver, and

etching to remove said porous silicon layer by immersing in a mixture of buffered hydrofluoric acid and an alcohol.

23. A method for preparing a semiconductor member, which comprises:

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forming a substrate having a non-porous silicon monocrystalline layer and a porous silicon layer, bonding another substrate having a surface made of an insulating material to the surface of said monocrystalline layer, and

etching to remove said porous silicon layer by immersing in a mixture of buffered hydrofluoric acid and hydrogen peroxide.

24. A method for preparing a semiconductor member, which comprises:

forming a substrate having a non-porous silicon monocrystalline layer and a porous silicon layer, bonding another substrate having a surface made of an insulating material to the surface of said monocrystalline layer, and

etching to remove said porous silicon layer by immersing in a mixture of buffered hydrofluoric acid, an alcohol and hydrogen peroxide.

5 25. A method for preparing a semiconductor member, which comprises the steps of: making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

bonding a light-transmissive glass substrate to the surface of said non-porous silicon monocrystalline layer, and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in hydrofluoric acid.

28. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on the silicon substrate made porous;

bonding another silicon substrate having an insulating layer on the surface thereof to the surface of said non-porous silicon monocrystalline layer; and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in hydrofluoric acid.

27. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous:

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

forming an oxide layer on the surface of said non-porous silicon monocrystalline layer:

bonding a light-transmissive substrate to the surface of said oxide layer, and

chemically etching to remove said silicon substrate made porous by immersing in hydrofluoric acid after coating the surfaces other than the surface of said silicon substrate made porous with a protecting material.

28. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous:

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

forming an oxide layer on the surface of said non-porous silicon monocrystalline layer,

bonding another silicon substrate having an insulating material on the surface thereof to the surface of said oxide layer on said non-porous silicon monocrystalline layer, and

chemically etching to remove said silicon substrate made porous by immersing in hydrofluoric acid.

29. A m thod for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous sillcon monocrystallin lay r n said silicon substrate med porous; bonding a light-transmissiv glass substrat t the surface of said non-porous silicon monocrystal-lin layer, and

s | ctively tching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate mad porous in a mixture of hydrofluoric acid and an alcohol.

30. A method for preparing a semiconductor member, which comprises the steps of:

making a sillcon substrate porous;

forming a non-porous silicon monocrystalline layer on the silicon substrate made porous;

bonding another silicon substrate having an insulating layer on the surface thereof to the surface of said non-porous silicon monocrystalline layer, and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in a mixture of hydrofluoric acid and an alcohol.

31. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

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forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

forming an oxide layer on the surface of said non-porous silicon monocrystalline layer;

bonding a light-transmissive substrate to the surface of said oxide layer, and

chemically etching to remove said silicon substrate made porous by immersing in a mixture of hydrotic acid and an alcohol after coating the surfaces other than the surface of said silicon substrate made porous with a protecting material.

32. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

forming an oxide layer on the surface of said non-porous silicon monocrystalline layer,

bonding another stilicon substrate having an insulating material on the surface thereof to the surface of said oxide layer on said non-porous silicon monocrystalline layer; and

chemically etching to remove said silicon substrate made porous by immersing in a mixture of hydrofluoric acid and an alcohol.

33. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

bonding a light-transmissive glass substrate to the surface of said non-porous silicon monocrystalline layer; and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing sald silicon substrate made porous in a mixture of hydrofluoric acid and hydrogen peroxide.

34. A method for preparing a semiconductor member, which comprises the steps of: making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on the silicon substrate made porous;

bonding another silicon substrate having an insulating layer on the surface thereof to the surface

of said non-porous silicon monocrystalline layer, and selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in a mixture of hydrofluonic acid and hydrogen peroxide.

35. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on sald silicon substrate made porous;

forming an oxide layer on the surface of said non-porous silicon monocrystalline layer;

bonding a light-transmissive substrate to the surface of said oxide layer, and

chemically etching to remove said silicon substrate made porous by immersing in a mixture comprising of hydrofluoric acid and hydrogen peroxide after coating the surfaces other than the surface of said silicon substrate made porous with a protecting material.

36. A m thod for preparing a s mic nductor m mber, which compris s th steps of:

making a silicon substrat porous;

forming a non-porous sillcon monocrystallin lay r on said sillcon substrate made porous;

forming an oxid layer on the surface of said non-porous silicon monocrystallin lay r,

bonding anoth relicon substrate having an insulating material in the surface thereof to the surface of eald oxid lay rion said non-porous silicon monocrystallin lay r, and

ch mically tching to remov sald silicon substrate mad porous by imm rsing in a mixtur of hydrofluoric acid and hydrogen p roxide.

37. A method for preparing a semiconductor member, which comprises the steps of:

making a sillcon substrate porous;

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forming a non-porous silicon monocrystalline layer on said allicon substrate made porous; bonding a light-transmissive glass substrate to the surface of said non-porous silicon monocrystalline layer, and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in a mixture of hydrofluoric acid, an alcohol and hydrogen peroxide.

38. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on the silicon substrate made porous;

bonding another silicon substrate having an insulating layer on the surface thereof to the surface of said non-porous silicon monocrystalline layer, and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in a mixture of hydrofluoric acid, an alcohol and hydrogen peroxide.

39. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

forming an oxide layer on the surface of said non-porous silicon monocrystalline layer;

bonding a light-transmissive substrate to the surface of said oxide layer, and

chemically etching to remove said silicon substrate made porous by immersing in a mixture of hydrofluoric acid, an alcohol and hydrogen peroxide after coating the surfaces other than the surface of said silicon substrate made prorous with a protecting material.

40. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

forming an oxide layer on the surface of said non-porous sillcon monocrystalline layer,

bonding another silicon substrate having an Insulating material on the surface thereof to the surface of said oxide layer on said non-porous silicon monocrystalline layer, and

chemically etching to remove said silicon substrate made porous by immersing in a mixture of hydrofluoric acid, an alcohol and hydrogen peroxide.

41. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

bonding a light-transmissive glass substrate to the surface of said non-porous silicon monocrystalline layer; and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in buffered hydrofluoric acid.

 A method for preparing a semiconductor member, which comprises the steps of: making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on the silicon substrate made porous;

bonding another silicon substrate having an insulating layer on the surface thereof to the surface of said non-porous silicon monocrystalline layer, and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in buffered hydrofluoric acid.

43. A method for pr paring a semiconductor member, which compris s th steps of:

making a silicon substrate porous;

forming a non-porous silic n monocrystalline lay r on said silicon substrate mad porous; forming an oxid layer n the surface of said non-porous silicon monocrystallin lay r;

bonding a light-transmissiv substrate to the surface of said oxide layer, and

chemically etching to remove said silicon substrat made porous by immersing in buffered hydroflucic acid after coating the surfaces ther than the surface of said silicon substrate mad porous with a protection material.

 A method for preparing a semiconductor member, which comprises the steps of: making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on sald silicon substrate made porous; forming an oxide layer on the surface of said non-porous silicon monocrystalline layer;

bonding another silicon substrate having an insulating material on the surface thereof to the surface of said oxide layer on said non-porous silicon monocrystalline layer; and

chemically etching to remove said sillcon substrate made porous by immersing in buffered hydrofluoric acid.

45. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous:

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous; bonding a light-transmissive glass substrate to the surface of said non-porous silicon monocrystalline layer; and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in a mixture of buffered hydrofluoric acid and an alcohol.

46. A method for preparing a semiconductor member, which comprises the steps of: making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on the silicon substrate made porous; bonding another silicon substrate having an insulating layer on the surface thereof to the surface

of said non-porous silicon monocrystalline layer, and selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in a mixture of buffered hydrofluoric acid and an alcohol.

47. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

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forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

forming an oxide layer on the surface of said non-porous silicon monocrystalline layer,

bonding a light-transmissive substrate to the surface of said oxide layer, and

chemically etching to remove said silicon substrate made porous by immersing in a mixture of buffered hydrofluoric acid and an alcohol after coating the surfaces other than the surface of said silicon substrate made porous with a protecting material.

48. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

forming an oxide layer on the surface of said non-porous silicon monocrystalline layer;

bonding another silicon substrate having an insulating material on the surface thereof to the surface of said oxide layer on said non-porous silicon monocrystalline layer; and

chemically etching to remove said silicon substrate made porous by immersing in a mixture of buffered hydrofluoric acid and an alcohol.

49. A method for preparing a semiconductor member, which comprises the steps of: making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

bonding a light-transmissive glass substrate to the surface of said non-porous silicon monocrystalline layer, and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in a mixture of buffered hydrofluoric acid and hydrogen peroxide.

50. A method for preparing a semiconductor member, which comprises the teps of:

making a silicon substrate porous;

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forming a non-porous silicon monocrystalline lay r on th licon substrate mad porous;

bonding another silicon substrate having an insulating lay ir on the surface thereof to the surface of said non-porous silicon monocrystalline lay ir, and

selectively etching porous silicon by chemically etching to remov porous silicon by immersing said silicon substrate made porous in a mixture of buffered hydrofluoric acid and hydrogen peroxide.

51. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous; forming a non-porous silicon monocrystalline layer on said silicon substrate made porous; forming an oxide layer on the surface of said non-porous silicon monocrystalline layer;

bonding a light-transmissive substrate to the surface of said oxide layer, and

chemically etching to remove said silicon substrate made porous by immersing in a mixture com-

chemically etching to remove said suiscan substrate made porous by immersing in a mixture comprising buffered hydrofluoric acid and hydrogen peroxide after coating the surfaces other than the surface of said silicon substrate made porous with a protecting material.

 A method for preparing a semiconductor member, which comprises the steps of: making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

forming an oxide layer on the surface of said non-porous sillcon monocrystalline layer,

bonding another silicon substrate having an insulating material on the surface thereof to the surface of said oxide layer on said non-porous silicon monocrystalline layer; and chemically etching to remove said silicon substrate made porous by immersing in a mixture of buf-

chemically exching to remove said silicon substrate made porous by immersing in a mixture of our fered hydrofluoric acid and hydrogen peroxide.

53. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said silicon substrate made porous;

bonding a light-transmissive glass substrate to the surface of said non-porous silicon monocrystalline layer, and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in a mixture of buffered hydrofluoric acid, an alcohol and hydrogen peroxide.

54. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on the silicon substrate made porous;

bonding another silicon substrate having an insulating layer on the surface thereof to the surface of said non-porous silicon monocrystalline layer, and

selectively etching porous silicon by chemically etching to remove porous silicon by immersing said silicon substrate made porous in a mixture of buffered hydrofluoric acid, an alcohol and hydrogen peroxide.

55. A method for preparing a semiconductor member, which comprises the steps of:

making a silicon substrate porous;

forming a non-porous silicon monocrystalline layer on said sillcon substrate made porous:

forming an oxide layer on the surface of said non-porous silicon monocrystalline layer;

bonding a light-transmissive substrate to the surface of said oxide layer, and

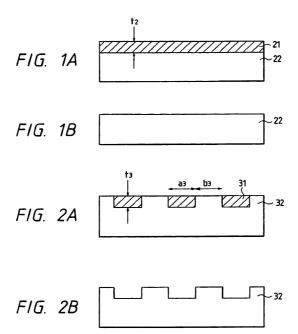
chemically etching to remove said silicon substrate made porous by immersing in a mixture of buffered hydrofluoric acid, an alcohol and hydrogen peroxide after coating the surfaces other than the surface of said silicon substrate made porous with a protecting material.

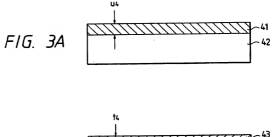
56. A method for preparing a semiconductor member, which comprises the steps of: making a silicon substrate porous;

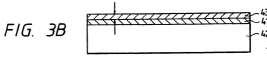
forming a non-porous silicon monocrystalline layer on said silicon substrate made porous; forming an oxide layer on the surface of said non-porous silicon monocrystalline layer:

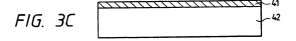
bonding another silicon substrate having an insulating material on the surface thereof to the surface of said oxide layer on said non-porous silicon monocrystalline layer; and

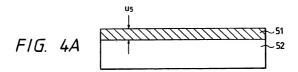
chemically etching to remove said silicon substrate made porous by immersing in a mixture comprising buffered hydrofluoric acid, an alcohol and hydrogen peroxide.

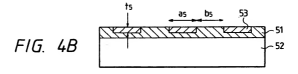


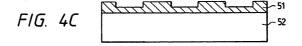


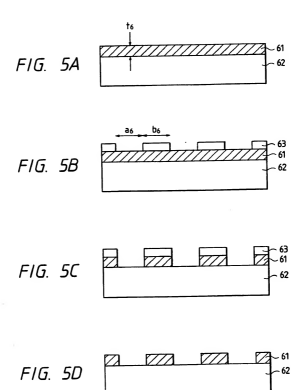


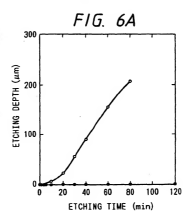


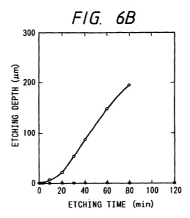


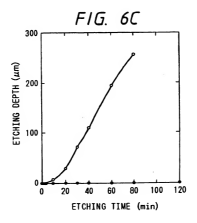


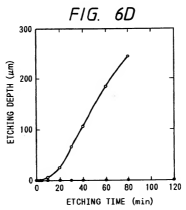


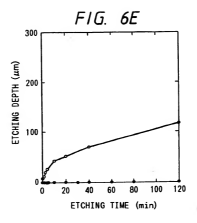


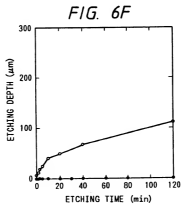


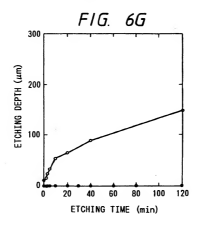


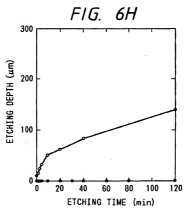


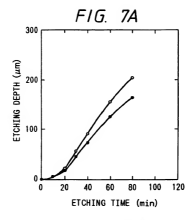


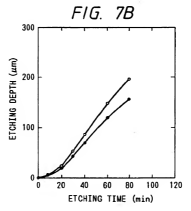


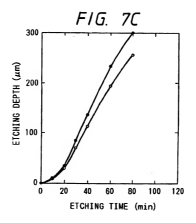


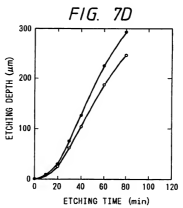


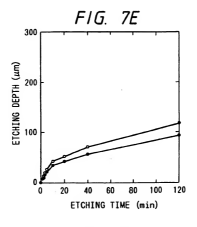


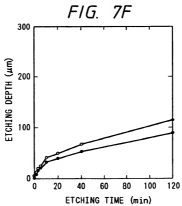


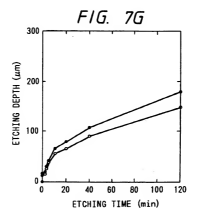


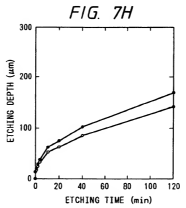


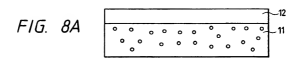


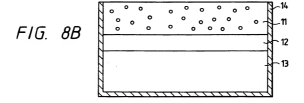


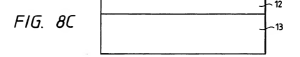


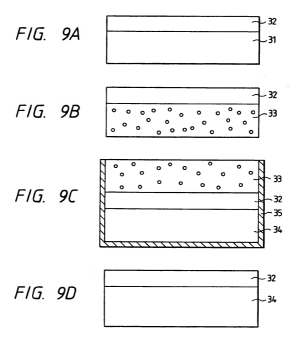


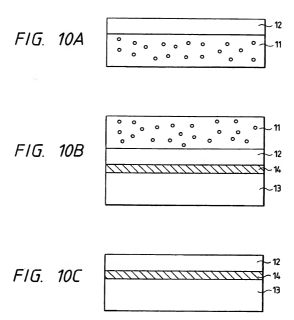


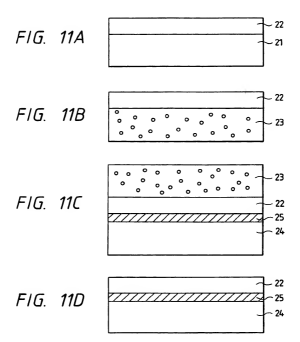


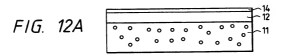


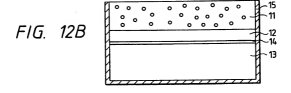


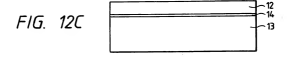


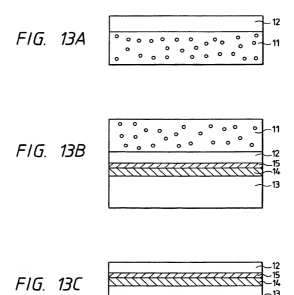


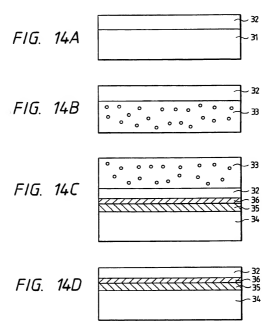
















## **EUROPEAN PATENT APPLICATION**

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(12)

(6) Int CL\*: H01L 21/306, H01L 21/20, H01L 21/76

(a) Priority: 15,02,91 JP 42212/91
12,02,91 JP 42212/91
12,02,91 JP 42212/91
12,02,91 JP 55504/91
12,02,91 JP 5550

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(7) Inventor: Sakaguchi, Klyofumi, oʻo Canon Kabushildi Kalsha 30-2, 3-chome, Shimomaruko Ohta-ku. Tokyo (JP) Inventor: Yonelara, Takao, oʻo Canon Kabushild Inventor: Yonelara, Takao, oʻo Canon Kabushild 30-2, 3-chome, Shimomaruko Ohta-ku. Tokyo (JP) Sababa Sato, Nobulniko, oʻo Canon Kabushild Kalsha Sato, Nobulniko, oʻo Canon Kabushild Kalsha Sato, Nobulniko, oʻo Canon Kabushild Ohta-ku. Tokyo (JP)

(74) Representative: Beresford, Keith Denis Lewis et all BERESFORD & Co. 2-5 Warnick Court High Holborn London WC1R 5DJ (GB)

- (43) Date of publication of application : 19,08.92 Bulletin 92/34
- (64) Designated Contracting States : AT BE CH DE DK ES FR GB GR IT LI LU NL PT SE
- (88) Date of deferred publication of search report : 01.03.95 Bulletin 95/09

(4) Etching solution for etching porous silicon, etching method using the etching solution and method of preparing semiconductor member using the etching solution.

A method for preparing a semiconductor member comprises:
 oming a substrate having a non-porous silicon monocrystalline layer and a porous silicon layer; monocrystalline layer and a porous silicon layer; a surface

made of an insulating material to the surface of the monocrystalline layer; and stching to remove the porous silicon layer by immersing in an etching solution.



# EUROPEAN SEARCH REPORT

EP 92 30 1252

	DOCUMENTS CONSID  Citation of document with indi				Relevant	CLASSIFICATION OF THE
Category	of relevant pass	Sea			claim	APPLICATION (Int.CLS)
x	APPLIED PHYSICS LETT vol.57, no.10, 3 Sep US		O, NEW YOR	RK 1,	9	H01L21/306 H01L21/20 H01L21/76
	pages 1046 - 1048 L.T. CANHAM 'SILCON FABRICATION BY ELECTI CHEMICAL DISSOLUTION	ROCHEMICAL OF WAFERS	AND			
	* page 1047, left co page 1048, left colu figure 3 *	nn, paragr	aph 2;			
>,χ	WO-A-91 09420 (SECR. HER BR. MAJEST. GOVE * claims 1,5,7; figu	RN. U.KG	FOR DEF.		,2,9,10	
٩	US-A-3 962 052 (IBM) * column 3, line 13	- line 36;	claim 1		, 17-56	
A	FR-A-2 374 396 (IBM)			1		
X	* claims 6-12 *			3	,5,7, 1,13,15	TECHNICAL FIELDS SEARCHED (Int.CL5)
A	IBM TECHNICAL DISCLO vol.14, no.11, April G.H. SCHWUTTKE ET AL WALLED PATTERNS IN (	1972, NEW ETCHING	YORK US OF VERTI	CAL		NOIL
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	pages 672 - 674 L. LIOU ET AL. 'AMOR BY ION IMPLANTATION. SOLUTIONS AND EFFECT	ETCHING R	LATE IN HE			
			-/			ļ
	The present search report has be	en drawn up for	all claims			
	Place of search		completion of the sec	ra l		Exeminer
	THE HAGUE	23 (	December	1994	Va	ncraeynest, F
X:2	CATEGORY OF CITED DOCUMENT articularly relevant if taken alone articularly relevant if combined with ano	nts	T : theory or E : earlier so	principle tent documents filing date	paderlying the	ne invention Mished on, or

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## EUROPEAN SEARCH REPORT

Application Number EP 92 30 1252

		DERED TO BE RELEVANT		
ategory	Citation of document with in of relevant pas		Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CL5)
1	JOURNAL OF THE ELECT vol.114, no.4, Apri' HAMPSHIRE US page 114	TROCHEMICAL SOCIETY, 1 1967, MANCHESTER, NEW		
	S.M. HU ET AL. 'OBSI	ERVATION OF ETCHING OF AQUEOUS HF SOLUTIONS'		
۸,	EP-A-0 469 630 (CANO * claims 1-125 *	ON)	17-56	
١.	EP-A-0 209 173 (PHI * claims 1-8,10,22,		17-56	# C
۸.	DEVICES AND MÁTERIA pages 33 - 36 A. NAKAGAWA ET AL. GATE BIPOLAR-MODE M	ERENCE ON SOLID-STATE LS 1988, 1988, TOKYO '500V LATERAL DOUBLE	17-56	
	DIRECT-BONDING (DIS * figure 1 *	DB)."		TECHNICAL FIELDS SEARCHED (In.Cl.5)
<b>A</b>	JOURNAL OF THE ELECTROCHEMICAL SOCIETY, vol. 135, no.8, August 1988, MANCHESTER, NEW HAMPSHIRE US pages 2105 - 2107 X Z. TU "FABRICATION OF SILICOON HICROSTRUCTURES BASED ON SELECTIVE FORMATION AND ETCHING OF POROUS SILICON" * page 2106, left column, paragraph 3 - right column, paragraph 8; figure 1 *			
	The present search report has			
	Place of search	Date of completing of the search		Domine
	THE HAGUE	23 December 199	ple underlying t	incraeynest, F
Y: 6	surticularly relevant if taken alone surticularly relevant if combined with as incursent of the same category	E : earlier patent d	date in the applicati	lon



# EUROPEAN SEARCH REPORT

EP 92 30 1252

	DOCUMENTS CONST	DERED TO BE RELEVA	N.I.			
ategory	Citation of document with in of relevant par		Relevant to claim	CLASSIFICATION OF THE APPLICATION (IntCLS)		
<b>A</b>	CHEM. PRUM., vol.27, no.12, 1977 pages 602 - 606 SEBEK SVATOPLUK 'CHI SURRACE OF SILICON' & CHEMICAL ABSTRACT vol.88, no.20, 15 M page 595 'ABSTRACT NO 145042	EMICAL ETCHING OF THE SINGLE CRYSTALS' S, ay 1978, OHIO	1,3,11			
<b>A</b>	PATENT ABSTRACTS OF vol. 14, no. 580 (E 1992 & JP-A-02 252 265 ( * abstract *		1-56			
				TECHNICAL FELDS SEARCHED (Int.Cl.5)		
	The present search report has i	ocn drawn up for all cinims  Dids of completes of the source		Dominio		
	THE HAGUE	23 December	1994 V	ancraeynest, F		
THE FIGURE  CATEGORY OF CITED DOCUMEN  X: particularly relevant if taken alone y: particularly relevant if constant with anot document of the same category A: technological background O: non-written disclosure P: intermediate document		NTS T: theory or pi E: earlier pate after the fi to ther D: document c L: document	rinciple underlying at document, but p ling date sized in the applica- ited for other reas	e underlying the invention sment, but published on, or te t the application		



EP 9 8 30 1252

	CLAIMS INCURRING FEES					
De.	The present European patent application comprised at the time of filling more than tan claims.					
(		All claims less have been paid within the prescribed time ilmit. The present Eurodean search report has been drawn up for all claims.				
1		Only oarn of the claims lees have been paid within the prescribed time limit. The present European search report has been grawn up for the first ten claims and for those claims for which claims fees have been paid.				
		namely claims:				
1		No ctaims (see nave been paid within the prescribed time limit. The present European acards report has been grawn up for the first len claims.				
		CLEAN DY DE DIE LEA CHE CHE DE				
	- LAC	CK OF UNITY OF INVENTION				
		Division considers that the present European petent application does not comply with the requirement of unity of				
inve nam		relates to several inventions or groups of inventions,				
ıem	ery:					
		See Sheet B.				
	Ø	All further search fees have been paid within the flaed time limit. The present European search report has been drawn up for all claims.				
	$\overline{}$	Only part of the further search fees have been paid within the fixed time limit. The present European search				
	Ц	report has been drawn up for those parts of the European petent application which relate to the inventions in respect of which search fees have been paid.				
		namely claims:				
		None of the further search fees has been paid within the fixed time first. The present European search record has been drawn up for those parts of the European patent application which relate to the invention first maniputed in the claims.				
		namely claims:				



EP-92301252

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# LACKS OF UNITY OF INVENTION

The Search Division considers that the present European patent application does not comply with the requirement of unity of inventions and relates to several inventions or proups of inventions.

- 1. Claims 1, 9: HF etchant for porous Si, and method for etching porous Si using HF etchant  $\,$
- 2. Claims 2-8, 10-16: Mixture of HF as etchant for porous Si, and method of etching using such a mixture of HF for porous Si
- 3. Claims 17-56: Formation of an SOI structure

6